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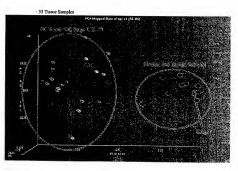
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54) Title: GENE EXPRESSION PROFILES IN BREAST TISSUE



(57) Abstract: The present invention results from the examination of tissue from breast carcinomas to identify genes differentially expressed between tumor biopsies and normal tissue. The invention includes diagnostic and screening methods using these genes as well as solid supports comprising oligonucleotide arrays that are complementary to or hybridize to the differentially expressed genes.

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GENE EXPRESSION PROFILES IN BREAST TISSUE

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RELATED APPLICATIONS

This application claims the priority of U.S. Provisional Application Nos. 60/263,757, filed January 25, 2001, 60/286,090, filed April 25, 2001, and 60/292,517, filed May 23, 2001, all of which are herein incorporated by reference in their entirety.

10 BACKGROUND OF THE INVENTION

One of the most pressing health issues today is breast cancer. In the industrial world, about one woman in every nine can expect to develop breast cancer in her lifetime. In the United States, it is the most common cancer amongst women, with an annual incidence of about 175,000 new cases and nearly 50,000 deaths. Despite an ongoing improvement in our understanding of the disease, breast cancer has remained resistant to medical intervention. Most clinical initiatives are focused on early diagnosis, followed by conventional forms of intervention, particularly surgery and chemotherapy. Such interventions are of limited success, particularly in patients where the tumor has undergone metastasis. There is a pressing need to improve the arsenal of therapies available to provide more precise and more effective treatment in a less invasive way. A promising area for the development of new modalities has emerged from recent understanding of the genetics of cancer.

One model used to characterize breast carcinogenesis asserts that normal cells undergo a multi-step process that broadly includes the steps of hyperplasia, pre-malignant change and in situ carcinoma. Multiple factors lead to atypical cell proliferation followed by carcinoma in situ. Carcinoma in situ is characterized as either ductal or lobular in form with the majority of invasive carcinomas being classified as ductal (85-95%). Among the ductal carcinomas, 15-20% encompass tubular, medullary, mucinous, papillary, adenoid, cystic, metaplastic, apocrine, squamous, secretory, lipid-rich, and cystic hypersecretory while the remaining ductal carcinomas are not specified.

To date, researchers have been able to identify a few genetic alterations believed to underlie tumor development. These genetic alterations include amplification of oncogenes and mutations that result in the loss of tumor suppressor genes. Tumor suppressor genes are genes that, in their wild-type alleles, express proteins that suppress abnormal cellular 2

proliferation. When the gene coding for a tumor suppressor protein is mutated or deleted, the resulting mutant protein or the complete lack of tumor suppressor protein expression may fail to correctly regulate cellular proliferation, and abnormal proliferation may take place, particularly if there is already existing damage to the cellular regulatory mechanism.

A number of well-studied human tumors and tumor cell lines have missing or non-functional tumor suppressor genes. Examples of tumor suppressor genes include, but are not limited to, the retinoblastoma susceptibility gene or RB gene, the p53 gene, the deletion in colon carcinoma (DCC) gene and the neurofibromatosis type 1 (NF-1) tumor suppressor gene (Weinberg, Science 254,1138-1146 (1991)). Loss of function or inactivation of tumor suppressor genes may play a central role in the initiation and/or progression of a significant number of human cancers.

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classification techniques.

Classification of heterogeneous populations of tumor types is a daunting task; yet, studies utilizing gene expression patterns to identify subtypes of cancer have produced initial results (see Perou, C. M. et al., Proc Natl Accad Sci USA 96, 9212-9217 (1999), Golub, T. R. et al., Science 286, 531-7 (1999), Alizadeh, A. A. et al., Nature 403, 503-11 (2000), Alon, U. et al. Proc Natl Acad Sci USA 96, 6745-50 (1999) and Bittner, M. et al., Nature 406, 536-40 (2000)). For example, molecular classification of B-cell lymphoma by gene expression profiling elucidated clinically distinct diffuse large-B-cell lymphoma butgroups (see Alizadeh supra). Stratification of patients based on their distinctive gene expression profiles may allow researchers to precisely group similar patient populations for evaluating chemotherapeutic agents. The more homogenous population of patients decreases the variability of patient-to-patient responses leading to the development of agents capable of eradicating specific subtypes of cancers previously unknown using standard

A study by Martin et al. (Cancer Res 60, 2232-8 (2000)) used a custom microarray composed of 124 genes discovered by differential display associated with either normal breast epithelial cells or from the MDA-MB-435 malignant breast tumor cell line. Using the custom microarray, researchers examined the relationship between expression patterns discovered by clustering a number of genes with clinical stages of breast cancer, indicating that gene expression patterns were capable of grouping breast tumors into distinct categories (Martin et al., supra).

The utilization of gene expression profiles to classify tumors, to identify drug targets, to identify diagnostic markers and/or to gain further insights into the consequences of chemotherapeutic treatments could facilitate the design of more efficacious patient—

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specific stratagems for treating a variety of cancers. In breast cancer, studies utilizing limited numbers of genes have classified tumors into subtypes based on gene expression profiles, and this study indicated a diversity of molecular phenotypes associated with breast tumors (Perou, C. M. et al., Nature 406, 747-52 (2000).

Although these studies have demonstrated that expression profiling may be used to produce improvements in diagnosis of breast cancer as well as the development of improved therapeutic strategies, further studies are needed as only a small portion of the genome was studied and analyses containing greater numbers of genes will advance our understanding of breast tumors even further. Accordingly, there remains a need in the art for materials and methods that permit a more accurate diagnosis of breast cancer and, in particular, ductal carcinoma. In addition, there remains a need in the art for methods to treat and methods to identify agents that can effectively treat breast cancer. The present invention meets these and other needs.

15 SUMMARY OF THE INVENTION

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The present invention is based on the discovery of the genes and their expression profiles associated with various types and stages of breast cancer.

The invention includes methods of diagnosing breast cancer in a patient comprising the step of detecting the level of expression in a tissue sample of two or more genes from Tables 1-5; wherein differential expression of the genes in Tables 1-5 is indicative of breast cancer.

The invention also includes methods of detecting the progression of breast cancer. For instance, methods of the invention include detecting the progression of breast cancer in a patient comprising the step of detecting the level of expression in a tissue sample of two or more genes from Tables 1-5; wherein differential expression of the genes in Tables 1-5 is indicative of breast cancer progression. In some preferred embodiments, PCA (Principal Component Analysis) based on all or a portion of the group of 50 genes identified in Table 1 may be used to differentiate between the different stages of breast cancer such as normal versus DCIS (ductal carcinoma in-situ) or DCIS versus microinvasive tissue samples. In some preferred embodiments, one or more genes may be selected from Tables 1, 3, 4 and/or 5.

In some aspects, the present invention provides a method of monitoring the treatment of a patient with breast cancer, comprising administering a pharmaceutical composition to the patient and preparing a gene expression profile from a cell or tissue

sample from the patient and comparing the patient gene expression profile to a gene expression from a cell population comprising normal breast cells or to a gene expression profile from a cell population comprising breast cancer cells or to both. In some preferred embodiments, the gene profile will include the expression level of one or more genes in Tables 1-5.

Another aspect of the present invention includes a method of treating a patient with breast cancer, comprising administering to the patient a pharmaceutical composition, wherein the composition alters the expression of at least one gene in Tables 1-5, preparing a gene expression profile from a cell or tissue sample from the patient comprising tumor cells and comparing the patient expression profile to a gene expression profile from an untreated cell population comprising breast cancer cells.

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In another aspect, the present invention provides a method of identifying ductal carcinoma in a patient, comprising detecting the level of expression in a tissue sample of two or more genes from Tables 1-5, wherein differential expression of the genes in Tables 1-5 is indicative of ductal carcinoma. In addition, by determining the expression level of two or more genes in the group of genes listed in Tables 1-5, one skilled in the art can differentiate between DCIS and a cribiform type of DCIS that is more prone to microinvasion.

In another aspect, the present invention provides a method of detecting the progression of carcinogenesis in a patient, comprising detecting the level of expression in a tissue sample of two or more genes from Tables 1-5; wherein differential expression of the genes in Tables 1-5 is indicative of breast carcinogenesis. Figures 6 and 7 are a graphical representation of how the genes listed in Table 5 cluster with disease stages in breast cancer.

The invention further includes methods of screening for an agent capable of modulating the onset or progression of breast cancer, comprising the steps of exposing a cell to the agent; and detecting the expression level of two or more genes from Tables 1-5. In some embodiments, the breast cancer may be a ductal carcinoma. In some preferred embodiments, one or more genes may be selected from a group consisting of those listed in Tables 1, 3, 4 and/or 5. In some preferred methods, it may be desirable to detect all or nearly all of the genes in the tables.

The invention further includes compositions comprising at least two oligonucleotides, wherein each of the oligonucleotides comprises a sequence that specifically hybridizes to a gene in Tables 1-5 as well as solid supports comprising at least two probes, wherein each of the probes comprises a sequence that specifically hybridizes to

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a gene in Tables 1-5. In some preferred embodiments, one or more genes may be selected from a group consisting of those listed in Tables 1, 3, 4 and/or 5.

The invention further includes computer systems comprising a database containing information identifying the expression level in breast tissue of a set of genes comprising at least two genes in Tables 1-5 and a user interface to view the information. In some preferred embodiments, one or more genes may be selected from a group consisting of those listed in Tables 1, 3, 4 and/or 5. The database may further include sequence information for the genes, information identifying the expression level for the set of genes in normal breast tissue and cancerous tissue and may contain links to external databases such as GenBank.

Lastly, the invention includes methods of using the databases, such as methods of using the disclosed computer systems to present information identifying the expression level in a tissue or cell of at least one gene in Tables 1-5, comprising the step of comparing the expression level of at least one gene in Tables 1-5 in the tissue or cell to the level of expression of the gene in the database. In some preferred embodiments, two or more genes may be selected from a group consisting of those listed in Tables 1, 3, 4 and/or 5.

BRIEF DESCRIPTION OF THE DRAWINGS

Figure 1 is an E-northern showing the expression of topoisomerase II alpha in various tissue types.

Figure 2 is an E-northern showing the expression of ICBP90 in various tissue types. Figure 3 is an E-northern showing the expression of MCT4 gene.

Figure 4 is an E-northern showing the expression of the frizzled related protein.

Figure 5 is an E-northern showing the expression of an EST Affy ID AI668620. Figure 6 is a PCA of the set of 28 samples using the top 50 genes identified by p-

values.

Figure 7 is a PCA of the set of 33 samples using the top 50 genes and ESTs identified by p-values.

Figure 8 is a PCA of the set of 91 samples using the top 31 myo-lamina genes and ESTs

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DETAILED DESCRIPTION OF THE PREFERRED EMBODIMENTS

Many biological functions are accomplished by altering the expression of various genes through transcriptional (e.g., through control of initiation, provision of RNA precursors, RNA processing, etc.) and/or translational control. For example, fundamental

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biological processes such as cell cycle, cell differentiation and cell death, are often characterized by the variations in the expression levels of groups of genes.

Changes in gene expression also are associated with pathogenesis. For example, the lack of sufficient expression of functional tumor suppressor genes and/or the over expression of oncogene/protooncogenes could lead to tumorgenesis or hyperplastic growth of cells (Marshall, Cell 64, 313-326 (1991); Weinberg, Science, 254, 1138-1146 (1991)). Thus, changes in the expression levels of particular genes (e.g., oncogenes or tumor suppressors) serve as signposts for the presence and progression of various diseases.

Monitoring changes in gene expression may also provide certain advantages during drug screening and development. Often drugs are pre-screened for the ability to interact with a major target without regard to other effects the drugs have on cells. Often such other effects cause toxicity in the whole animal, which prevent the development and use of the potential drug.

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Applicants have examined samples from normal breast tissue and from cancerous breast tissue to identify global changes in gene expression between tumor biopsies and normal tissue. These global changes in gene expression, also referred to as expression profiles, provide useful markers for diagnostic uses as well as markers that can be used to monitor disease states, disease progression, drug toxicity, drug efficacy and drug metabolism

The gene expression profiles described herein were derived from normal and tumor samples from female patients between the ages of 39 and 52 years old, and were from three different ethnic origins (Caucasian, African-American and Asian). Infiltrating Ductal Carcinoma (IDC) patient samples were studied for cancer-related expression, as 85% of the breast cancer patients were afflicted with this form of the disease.

Histological analysis of each tissue sample was performed and samples were segregated into either normal or malignant categories. The normal tissue samples were acquired from neighboring tissue of patients suffering from one of the following disorders: macromastia, mild fibrosis, infiltrating lobular carcinoma, or infiltrating ducal carcinoma, however; each tissue was diagnosed as normal by histological analysis. Samples were also characterized by the type and grade of IDC for each patient sample utilized in the study.

The present invention provides compositions and methods to detect the level of expression of genes that may be differentially expressed dependent upon the state of the cell, i.e., normal versus cancerous. These expression profiles of genes provide molecular tools for evaluating toxicity, drug efficacy, drug metabolism, development, and disease

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monitoring. Changes in the expression profile from a baseline profile can be used as an indication of such effects. Those skilled in the art can use any of a variety of known techniques to evaluate the expression of one or more of the genes and/or gene fragments identified in the instant application in order to observe changes in the expression profile in a tissue or sample of interest.

Definitions

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In the description that follows, numerous terms and phrases known to those skilled in the art are used. In the interest of clarity and consistency of interpretation, the definitions of certain terms and phrases are provided.

As used herein, the phrase "detecting the level of expression" includes methods that quantify expression levels as well as methods that determine whether a gene of interest is expressed at all. Thus, an assay which provides a yes or no result without necessarily providing quantification of an amount of expression is an assay that requires "detecting the level of expression" as that phrase is used herein.

As used herein, oligonucleotide sequences that are complementary to one or more of the genes described herein, refers to oligonucleotides that are capable of hybridizing under stringent conditions to at least part of the nucleotide sequence of said genes. Such hybridizable oligonucleotides will typically exhibit at least about 75% sequence identity at the nucleotide level to said genes, preferably about 80% or 85% sequence identity or more preferably about 90% or 95% or more nucleotide sequence identity to said genes.

"Bind(s) substantially" refers to complementary hybridization between a probe nucleic acid and a target nucleic acid and embraces minor mismatches that can be accommodated by reducing the stringency of the hybridization media to achieve the desired detection of the target polynucleotide sequence.

The terms "background" or "background signal intensity" refer to hybridization signals resulting from non-specific binding, or other interactions, between the labeled target nucleic acids and components of the oligonucleotide array (e.g., the oligonucleotide probes, control probes, the array substrate, etc.). Background signals may also be produced by intrinsic fluorescence of the array components themselves. A single background signal can be calculated for the entire array, or a different background signal may be calculated for each target nucleic acid. In a preferred embodiment, background is calculated as the average hybridization signal intensity for the lowest 5% to 10% of the probes in the array, or, where a different background signal is calculated for each target gene, for the lowest 5%

to 10% of the probes for each gene. Of course, one of skill in the art will appreciate that where the probes to a particular gene hybridize well and thus appear to be specifically binding to a target sequence, they should not be used in a background signal calculation. Alternatively, background may be calculated as the average hybridization signal intensity produced by hybridization to probes that are not complementary to any sequence found in the sample (e.g., probes directed to nucleic acids of the opposite sense or to genes not found in the sample such as bacterial genes where the sample is mammalian nucleic acids). Background can also be calculated as the average signal intensity produced by regions of the array that lack any probes at all.

The phrase "hybridizing specifically to" refers to the binding, duplexing or hybridizing of a molecule substantially to or only to a particular nucleotide sequence or sequences under stringent conditions when that sequence is present in a complex mixture (e.g., total cellular) DNA or RNA.

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Assays and methods of the invention may utilize available formats to simultaneously screen at least about 100, preferably about 1000, more preferably about 10,000 and most preferably about 1,000,000 or more different nucleic acid hybridizations.

The terms "mismatch control" or "mismatch probe" refer to a probe whose sequence is deliberately selected not to be perfectly complementary to a particular target sequence. For each mismatch (MM) control in a high-density array there typically exists a corresponding perfect match (PM) probe that is perfectly complementary to the same particular target sequence. The mismatch may comprise one or more bases that are not complementary to the corresponding bases of the target sequence.

While the mismatch(s) may be located anywhere in the mismatch probe, terminal mismatches are less desirable as a terminal mismatch is less likely to prevent hybridization of the target sequence. In a particularly preferred embodiment, the mismatch is located at or near the center of the probe such that the mismatch is most likely to destabilize the duplex with the target sequence under the test hybridization conditions.

The term "perfect match probe" refers to a probe that has a sequence that is perfectly complementary to a particular target sequence. The test probe is typically perfectly complementary to a portion (subsequence) of the target sequence. The perfect match (PM) probe can be a "test probe", a "normalization control" probe, an expression level control . probe and the like. A perfect match control or perfect match probe is, however, distinguished from a "mismatch control" or "mismatch probe."

As used herein a "probe" is defined as a nucleic acid, preferably an oligonucleotide, capable of binding to a target nucleic acid of complementary sequence through one or more types of chemical bonds, usually through complementary base pairing, usually through hydrogen bond formation. As used herein, a probe may include natural (i.e., A, G, U, C or T) or modified bases (7-deazaguanosine, inosine, etc.). In addition, the bases in probes may be joined by a linkage other than a phosphodiester bond, so long as it does not interfere with hybridization. Thus, probes may be peptide nucleic acids in which the constituent bases are joined by peptide bonds rather than phosphodiester linkages.

The term "stringent conditions" refers to conditions under which a probe will hybridize to its target subsequence, but with only insubstantial hybridization to other sequences or to other sequences such that the difference may be identified. Stringent conditions are sequence-dependent and will be different in different circumstances. Longer sequences hybridize specifically at higher temperatures. Generally, stringent conditions are selected to be about 5°C lower than the thermal melting point (Tm) for the specific sequence at a defined ionic strength and off.

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Typically, stringent conditions will be those in which the salt concentration is at least about 0.01 to 1.0 M sodium ion concentration (or other salts) at pH 7.0 to 8.3 and the temperature is at least about 30°C for short probes (e.g., 10 to 50 nucleotide). Stringent conditions may also be achieved with the addition of destabilizing agents such as formamide.

The "percentage of sequence identity" or "sequence identity" is determined by comparing two optimally aligned sequences or subsequences over a comparison window or span, wherein the portion of the polynucleotide sequence in the comparison window may optionally comprise additions or deletions (i.e., gaps) as compared to the reference sequence (which does not comprise additions or deletions) for optimal alignment of the two sequences. The percentage is calculated by determining the number of positions at which the identical subunit (e.g., nucleic acid base or amino acid residue) occurs in both sequences to yield the number of matched positions, dividing the number of matched positions by the total number of positions in the window of comparison and multiplying the result by 100 to yield the percentage of sequence identity. Percentage sequence identity when calculated using the programs GAP or BESTFIT (see below) is calculated using default gap weights.

Homology or identity may be determined by BLAST (Basic Local Alignment Search Tool) analysis using the algorithm employed by the programs blastp, blastn, blastx,

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tblastn and tblastx (Karlin et al., Proc Natl Acad Sci USA 87, 2264-2268 (1990) and Altschul, J Mol Evol 36, 290-300 (1993), fully incorporated by reference) which are tailored for sequence similarity searching. The approach used by the BLAST program is to first consider similar segments between a query sequence and a database sequence, then to evaluate the statistical significance of all matches that are identified and finally to summarize only those matches which satisfy a preselected threshold of significance. For a discussion of basic issues in similarity searching of sequence databases, see Altschul et al., (Nature Genet 6, 119-129 (1994)) which is fully incorporated by reference. The search parameters for histogram, descriptions, alignments, expect (i.e., the statistical significance threshold for reporting matches against database sequences), cutoff, matrix and filter are at the default settings. The default scoring matrix used by blastp, blastx, tblastn, and tblastx is the BLOSUM62 matrix (Henikoff et al., Proc Natl Acad Sci USA 89, 10915-10919, (1992) fully incorporated by reference). Four blastn parameters were adjusted as follows: O=10 (gap creation penalty); R=10 (gap extension penalty); wink=1 (generates word hits at every winkth position along the query); and gapw=16 (sets the window width within which gapped alignments are generated). The equivalent Blastp parameter settings were O=9; R=2; wink=1; and gapw=32. A Bestfit comparison between sequences, available in the GCG package version 10.0, uses DNA parameters GAP=50 (gap creation penalty) and LEN=3 (gap extension penalty) and the equivalent settings in protein comparisons are GAP=8 and

Uses of Differentially Expressed Genes

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The present invention identifies those genes differentially expressed between normal breast tissue and cancerous breast tissue. One of skill in the art can select one or more of the genes identified as being differentially expressed in Tables 1-5 and use the information and methods provided herein to interrogate or test a particular sample. For a particular interrogation of two conditions or sources, it may be desirable to select those genes which display a great deal of difference in the expression pattern between the two conditions or sources. At least a two-fold difference may be desirable, but a three-fold, five-fold or ten-fold difference may be preferred in some instances. Interrogations of the genes or proteins can be performed to yield different information.

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As described herein, the genes and gene expression information provided in Tables 1-5 may be used as diagnostic markers for the prediction or identification of the malignant state of breast tissue. For instance, a breast tissue sample or other sample from a patient may be assayed by any of the methods known to those skilled in the art, and the expression levels from one or more genes from Tables 1-5, may be compared to the expression levels found in normal breast tissue, tissue from breast carcinoma or both. Expression profiles generated from the tissue or other samples that substantially resemble an expression profile from normal or diseased breast tissue may be used, for instance, to aid in disease diagnosis. Comparison of the expression data, as well as available sequence or other information may be done by researcher or diagnostician or may be done with the aid of a computer and databases as described herein.

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For example, genes over-expressed by 3-fold or greater, as well as having the smallest p-values from a t-test, were discovered by comparing 13 normal tissue samples and 15 infiltrating ductal carcinoma tissue samples composed of mostly stage II and III tissue samples. This analysis provided a set of genes (listed in Table 1) capable of distinguishing between the 13 normal and 15 tumor samples by PCA (Principal Component Analysis). In order to evaluate the ability of the genes to distinguish between normal and tumor tissue samples, a group of 33 tissues was selected from an existing gene expression database composed of normal, benign, DCIS (ductal carcinoma in-situ), microinvasive, stage I, stage 20 II. and stage III breast cancer samples. PCA of the 33 tissue samples indicated that the genes selected based on the smallest p-values classified 32 out of 33 tissue samples correctly, while one stage I tissue sample was misclassified as a normal sample. Accordingly, these genes can be used diagnostically to differentiate normal/benign samples from tissue samples containing intraductal or infiltrating ductal carcinoma of the breast.

In another study, the PCA based on this group of genes indicates that these genes may be used to differentiate between the different stages of breast cancer such as normal versus DCIS or DCIS versus microinvasive tissue samples as graphically shown in Figures 6 and 7. The DCIS sample that contained focal microinvasions was grouped with the Stage I and II tumor samples. This group of genes may be used to determine if a DCIS sample contains microinvasions.

Use of the Breast Cancer Markers for Monitoring Disease Progression

Molecular expression markers for breast cancer can be used to confirm the type and progression of cancer made on the basis of morphological criteria. For example, normal

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breast tissue could be distinguished from invasive carcinoma based on the level and type of genes expressed in a tissue sample. In some situations, identifications of cell type or source is ambiguous based on classical criteria. In these situations, the molecular expression markers of the present invention are useful.

In addition, progression of ductal carcinoma in situ to microinvasive carcinoma can be monitored by following the expression patterns of the involved genes using the molecular expression markers of the present invention. Monitoring of the efficacy of certain drug regimens can also be accomplished by following the expression patterns of the molecular expression markers.

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In addition to the different disease progression stages which have been shown in Figures 6-7, as shown in the examples below, other developmental stages can be identified using these same molecular expression markers. While the importance of these markers in development has been shown here, variations in their expression may occur at other times. For example, variation in the expression level of one or more of the marker genes identified herein may be use to distinguish benign stages of breast cancer from malignant states.

As described above, the genes and gene expression information provided in Tables 1-5 may also be used as markers for the direct monitoring of disease progression, for instance, the development of breast cancer. For instance, a breast tissue sample or other sample from a patient may be assayed by any of the methods known to those of skill in the art, and the expression levels in the sample from a gene or genes from Tables 1-5 may be compared to the expression levels found in normal breast tissue, tissue from breast cancer or both. Comparison of the expression data, as well as available sequence or other information may be done by researcher or diagnostician or may be done with the aid of a computer and databases as described herein.

For instance, methods of this invention may use the 35 gene group (profile) composed of genes expressed in myoepithelial cells and basal lamina components in Table 3. The absence of both myoepithelial cells or basement membrane components usually indicates that the intraductal carcinoma is invasive. This group of 35 genes listed in Table 3 may be used to determine if myoepithelial and/or basal lamina components are present in a tissue sample. It includes 23 genes exhibiting a fold change of 3 fold or higher and 12 genes displaying a change of less than 3 fold. This group of 23 genes was used to distinguish between normal and tumor samples for a group of 33 tissue samples. In this study, the 23 genes were able to classify 32 out of 33 samples correctly and 26 out of 28 samples used to isolate this subgroup of genes. This group of genes can be used to identify

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the various stages of ductal carcinoma tissues more discretely than the 50-gene set. The study also demonstrates that this group of genes can differentiate between DCIS and a cribiform type of DCIS that is more prone to microinvasion. Clinically, the ability to discern DCIS with microinvasions or phenotypes prone to microinvasions such as the cribiform type would allow subgrouping of the samples containing microinvasions as a type of patient that should be treated more aggressively than DCIS patients lacking this gene expression fingerprint. A subclass of DCIS (cribiform type) based on the gene expression fingerprint may be subgrouped as a micro invasive sample based on the gene expression pattern associated with this sample.

Use of the Breast Cancer Markers for Drug Screening

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According to the present invention, potential drugs can be screened to determine if application of the drug alters the expression of one or more of the genes identified herein. This may be useful, for example, in determining whether a particular drug is effective in treating a particular patient with breast cancer. In the case where a gene's expression is affected by the potential drug such that its level of expression returns to normal, the drug is indicated in the treatment of breast cancer. Similarly, a drug which causes expression of a gene which is not normally expressed by epithelial cells in the breast, may be contraindicated in the treatment of breast cancer.

According to the present invention, the genes identified in Tables 1-5 may also be used as markers to evaluate the effects of a candidate drug or agent on a cell, particularly a cell undergoing malignant transformation, for instance, a breast cancer cell or tissue sample. A candidate drug or agent can be screened for the ability to stimulate the transcription or expression of a given marker or markers (drug targets) or to down-regulate or inhibit the transcription or expression of a marker or markers. According to the present invention, one can also compare the specificity of a drug's effects by looking at the number of markers affected by the drug and comparing them to the number of markers affected by a different drug. A more specific drug will affect fewer transcriptional targets. Similar sets of markers identified for two drugs indicates a similarity of effects.

Assays to monitor the expression of a marker or markers as defined in Tables 1-5 may utilize any available means of monitoring for changes in the expression level of the nucleic acids of the invention. As used herein, an agent is said to modulate the expression of a nucleic acid of the invention if it is capable of up- or down-regulating expression of the nucleic acid in a cell.

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Agents that are assayed in the above methods can be randomly selected or rationally selected or designed. As used herein, an agent is said to be randomly selected when the agent is chosen randomly without considering the specific sequences involved in the association of the a protein of the invention alone or with its associated substrates, binding partners, etc. An example of randomly selected agents is the use a chemical library or a pentide combinatorial library, or a growth broth of an organism.

As used herein, an agent is said to be rationally selected or designed when the agent is chosen on a nonrandom basis which takes into account the sequence of the target site and/or its conformation in connection with the agents action. Agents can be selected or designed by utilizing the peptide sequences that make up these sites. For example, a rationally selected peptide agent can be a peptide whose amino acid sequence is identical to or a derivative of any functional consensus site.

The agents of the present invention can be, as examples, peptides, small chemical molecules, vitamin derivatives, as well as carbohydrates, lipids, oligonucleotides and covalent and non-covalent combinations thereof. Dominant negative proteins, DNA encoding these proteins, antibodies to these proteins, peptide fragments of these proteins or mimics of these proteins may be introduced into cells to affect function. "Mimic" as used herein refers to the modification of a region or several regions of a peptide molecule to provide a structure chemically different from the parent peptide but topographically and functionally similar to the parent peptide (see Grant in Molecular Biology and Biotechnology, Meyers, ed., VCH Publishers (1995)). A skilled artisan can readily recognize that there is no limit as to the structural nature of the agents of the present invention.

25 Assav Formats

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The genes identified as being differentially expressed in breast cancer may be used in a variety of nucleic acid detection assays to detect or quantify the expression level of a gene or multiple genes in a given sample. For example, traditional Northern blotting, nuclease protection, RT-PCR and differential display methods may be used for detecting gene expression levels.

The protein products of the genes identified herein can also be assayed to determine the amount of expression. Methods for assaying for a protein include Western blot, immunoprecipitation, radioimmunoassay. It is preferred, however, that the mRNA be assayed as an indication of expression. Methods for assaying for mRNA include Northern

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blots, slot blots, dot blots, and hybridization to an ordered array of oligonucleotides. Any method for specifically and quantitatively measuring a specific protein or mRNA or DNA product can be used. However, methods and assays of the invention are most efficiently designed with PCR or array or chip hybridization-based methods for detecting the expression of a large number of genes.

Any hybridization assay format may be used, including solution-based and solid support-based assay formats. A preferred solid support is a high density array also known as a DNA chip or a gene chip. One variation of the DNA chip contains hundreds of thousands of discrete microscopic channels that pass completely through it. Probe molecules are attached to the inner surface of these channels, and molecules from the samples to be tested flow through the channels, coming into close proximity with the probes for hybridization. In one assay format, gene chips containing probes to at least two genes from Tables 1-5 may be used to directly monitor or detect changes in gene expression in the treated or exposed cell as described herein. Assays of the invention may measure the expression levels of about one, two, three, five, seven, ten, 15, 20, 25, 50, 100 or more genes in the Tables.

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The genes and ESTs of the present invention may be assayed in any convenient sample form. For example, samples may be assayed in the form mRNA or reverse transcribed mRNA. Samples may be cloned or not and the samples or individual genes may be amplified or not. The cloning itself does not appear to bias the representation of genes within a population. However, it may be preferable to use polyA+RNA as a source, as it can be used with less processing steps. In some embodiments, it may be preferable to assay the protein or pertide expressed by the gene.

The sequences of the expression marker genes of Tables 1-5 are available in the public databases. Tables 1-5 provide the Accession numbers and name for each of the sequences. The sequences of the genes in GenBank are herein expressly incorporated by reference in their entirety as of the filing date of this application. (see www.ncbi.nim.nih.gov).

Additional assay formats may be used to monitor the ability of the agent to modulate the expression of a gene identified in Tables 1-5. For instance, as described above, mRNA expression may be monitored directly by hybridization of probes to the nucleic acids of the invention. Cell lines are exposed to an agent to be tested under appropriate conditions and time and total RNA or mRNA is isolated by standard procedures such those disclosed in Sambrook et al., Molecular Cloning - A Laboratory Manual, Cold Spring Harbor

Laboratory Press, Cold Spring Harbor, NY (1989)). In some embodiments, it may be desirable to amplify one or more of the RNA molecules isolated prior to application of the RNA to the gene chip. Using techniques well known in the art, the RNA may be reverse transcribed and amplified in the form of DNA or may be reverse transcribed into DNA and the DNA used as a template for transcription to generate recombinant RNA. Any method that results in the production of a sufficient quantity of nucleic acid to be hybridized effectively to the gene chip may be used.

In another format, cell lines that contain reporter gene fusions between the open reading frame and or the 3° or 5° regulatory regions of a gene in Tables 1-5 and any assayable fusion partner may be prepared. Numerous assayable fusion partners are known and readily available including the firefly luciferase gene and the gene encoding chloramphenicol acetyltransferase (Alam et al., Anal Biochem 188, 245-254 (1990)). Cell lines containing the reporter gene fusions are then exposed to the agent to be tested under appropriate conditions and time. Differential expression of the reporter gene between samples exposed to the agent and control samples identifies agents which modulate the expression of the nucleic acid.

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In another assay format, cells or cell lines are first identified which express one or more of the gene products of the invention physiologically. Cells and/or cell lines so identified would preferably comprise the necessary cellular machinery to ensure that the transcriptional and/or translational apparatus of the cells would faithfully mimic the response of normal or cancerous breast tissue to an exogenous agent. Such machinery would likely include appropriate surface transduction mechanisms and/or cytosolic factors. Such cell lines may be, but are not required to be, derived from breast tissue. The cells and/or cell lines may then be contacted with an agent and the expression of one or more of the genes of interest may then be assayed. The genes may be assayed at the mRNA level and/or at the protein level.

In some embodiments, such cells or cell lines may be transduced or transfected with an expression vehicle (e.g., a plasmid or viral vector) containing an expression construct comprising an operable 5'-promoter containing end of a gene of interest identified in Tables 1-5 fused to one or more nucleic acid sequences encoding one or more antigenic fragments. The construct may comprise all or a portion of the coding sequence of the gene of interest which may be positioned 5'- or 3'- to a sequence encoding an antigenic fragment. The coding sequence of the gene of interest may be translated or un-translated after transcription of the gene fusion. At least one antigenic fragment may be translated. The antigenic

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fragments are selected so that the fragments are under the transcriptional control of the promoter of the gene of interest and are expressed in a fashion substantially similar to the expression pattern of the gene of interest. The antigenic fragments may be expressed as polypeptides whose molecular weight can be distinguished from the naturally occurring polypeptides. In some embodiments, gene products of the invention may further comprise an immunologically distinct tag. Such a process is well known in the art (see Sambrook et al., supra).

Cells or cell lines transduced or transfected as outlined above are then contacted with agents under appropriate conditions; for example, the agent comprises a pharmaceutically acceptable excipient and is contacted with cells comprised in an aqueous physiological buffer such as phosphate buffered saline (PBS) at physiological pH, Eagles balanced salt solution (BSS) at physiological pH, PBS or BSS comprising serum or conditioned media comprising PBS or BSS and serum incubated at 37°C. Said conditions may be modulated as deemed necessary by one of skill in the art. Subsequent to contacting the cells with the agent, said cells will be disrupted and the polypeptides of the lysate are fractionated such that a polypeptide fraction is pooled and contacted with an antibody to be further processed by immunological assay (e.g., ELISA, immunoprecipitation or Western blot). The pool of proteins isolated from the "agent-contacted" sample will be compared with a control sample where only the excipient is contacted with the cells and an increase or decrease in the immunologically generated signal from the "agent-contacted" sample compared to the control will be used to distinguish the effectiveness of the agent.

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Another embodiment of the present invention provides methods for identifying agents that modulate the levels, concentration or at least one activity of a protein(s) encoded by the genes in Tables 1-5. Such methods or assays may utilize any means of monitoring or detecting the desired activity.

In one format, the relative amounts of a protein of the invention produced in a cell population that has been exposed to the agent to be tested may be compared to the amount produced in an un-exposed control cell population. In this format, probes such as specific antibodies are used to monitor the differential expression of the protein in the different cell populations. Cell lines or populations are exposed to the agent to be tested under appropriate conditions and time. Cellular lysates may be prepared from the exposed cell line or population and a control, unexposed cell line or population. The cellular lysates are then analyzed with the probe, such as a specific antibody.

Probe Design

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Probes based on the sequences of the genes described herein may be prepared by any commonly available method. Oligonucleotide probes for assaying the tissue or cell sample are preferably of sufficient length to specifically hybridize only to appropriate, complementary genes or transcripts. Typically the oligonucleotide probes will be at least 10, 12, 14, 16, 18, 20 or 25 nucleotides in length. In some cases longer probes of at least 30, 40, or 50 nucleotides will be desirable.

One of skill in the art will appreciate that an enormous number of array designs are suitable for the practice of this invention. The high density array will typically include a number of probes that specifically hybridize to the sequences of interest. See WO 99/32660 for methods of producing probes for a given gene or genes. In addition, in a preferred embodiment, the array will include one or more control probes.

High density array chips of the invention include "test probes." Test probes may be oligonucleotides that range from about 5 to about 500 or about 50 moleotides, more preferably from about 10 to about 40 nucleotides and most preferably from about 15 to about 40 nucleotides in length. In other particularly preferred embodiments, the probes are about 20 or 25 nucleotides in length. In another preferred embodiment, test probes are double or single strand DNA sequences. DNA sequences may be isolated or cloned from natural sources or amplified from natural sources using natural nucleic acid as templates. These probes have sequences complementary to particular subsequences of the genes whose expression they are designed to detect. Thus, the test probes are capable of specifically hybridizing to the target nucleic acid they are to detect.

In addition to test probes that bind the target nucleic acid(s) of interest, the high density array can contain a number of control probes. The control probes fall into three categories referred to herein as (1) normalization controls; (2) expression level controls; and (3) mismatch controls.

Normalization controls are oligonucleotide or other nucleic acid probes that are complementary to labeled reference oligonucleotides or other nucleic acid sequences that are added to the nucleic acid sample. The signals obtained from the normalization controls after hybridization provide a control for variations in hybridization conditions, label intensity, "reading" efficiency and other factors that may cause the signal of a perfect hybridization to vary between arrays. In a preferred embodiment, signals (e.g., fluorescence

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intensity) read from all other probes in the array are divided by the signal (e.g., fluorescence intensity) from the control probes thereby normalizing the measurements.

Virtually any probe may serve as a normalization control. However, it is recognized that hybridization efficiency varies with base composition and probe length. Preferred normalization probes are selected to reflect the average length of the other probes present in the array, however, they can be selected to cover a range of lengths. The normalization control(s) can also be selected to reflect the (average) base composition of the other probes in the array, however in a preferred embodiment, only one or a few probes are used and they are selected such that they hybridize well (i.e., no secondary structure) and do not match any target-specific probes.

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Expression level controls are probes that hybridize specifically with constitutively expressed genes in the biological sample. Virtually any constitutively expressed gene provides a suitable target for expression level controls. Typical expression level control probes have sequences complementary to subsequences of constitutively expressed "housekeeping genes" including, but not limited to the β-actin gene, the transferrin receptor gene, the GAPDH gene, and the like.

Mismatch controls may also be provided for the probes to the target genes, for expression level controls or for normalization controls. Mismatch controls are oligonucleotide probes or other nucleic acid probes identical to their corresponding test or control probes except for the presence of one or more mismatched bases. A mismatched base is a base selected so that it is not complementary to the corresponding base in the target sequence to which the probe would otherwise specifically hybridize. One or more mismatches are selected such that under appropriate hybridization conditions (e.g., stringent conditions) the test or control probe would be expected to hybridize with its target sequence, but the mismatch probe would not hybridize (or would hybridize to a significantly lesser extent). Preferred mismatch probes contain a central mismatch. Thus, for example, where a probe is a twenty-mer, a corresponding mismatch probe may have the identical sequence except for a single base mismatch (e.g., substituting a G, a C or a T for an A) at any of positions 6 through 14 (the central mismatch).

Mismatch probes thus provide a control for non-specific binding or cross hybridization to a nucleic acid in the sample other than the target to which the probe is directed. Mismatch probes also indicate whether a hybridization is specific or not. For example, if the target is present the perfect match probes should be consistently brighter

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than the mismatch probes. In addition, if all central mismatches are present, the mismatch probes can be used to detect a mutation. The difference in intensity between the perfect match and the mismatch probe (I(PM) - I(MM)) provides a good measure of the concentration of the hybridized material.

Nucleic Acid Samples

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As is apparent to one of ordinary skill in the art, nucleic acid samples used in the methods and assays of the invention may be prepared by any available method or process. Methods of isolating total mRNA are also well known to those of skill in the art. For example, methods of isolation and purification of nucleic acids are described in detail in Chapter 3 of <u>Laboratory Techniques in Biochemistry and Molecular Biology, Vol. 24</u>, Hybridization With Nucleic Acid Probes: Theory and Nucleic Acid Probes, P. Tijssen, ed., Elsevier Press, New York (1993). Such samples include RNA samples, but also include cDNA synthesized from a mRNA sample isolated from a cell or tissue of interest. Such samples also include DNA amplified from the cDNA, and an RNA transcribed from the amplified DNA. One of skill in the art would appreciate that it may be desirable to inhibit or destroy RNase present in homogenates before homogenates can be used.

Biological samples may be of any biological tissue or fluid or cells from any organism as well as cells raised in vitro, such as cell lines and tissue culture cells.

Frequently the sample will be a "clinical sample" which is a sample derived from a patient. Typical clinical samples include, but are not limited to, breast tissue biopsy, sputum, blood, blood-cells (e.g., white cells), tissue or fine needle biopsy samples, urine, peritoneal fluid, and pleural fluid, or cells therefrom.

Biological samples may also include sections of tissues, such as frozen sections or formalin fixed sections taken for histological purposes.

Solid Supports

Solid supports containing oligonucleotide probes for differentially expressed genes can be any solid or semisolid support material known to those skilled in the art. Suitable examples include, but are not limited to, membranes, filters, tissue culture dishes, polyvinyl chloride dishes, beads, test strips, silicon or glass based chips and the like. Suitable glass wafers and hybridization methods are widely available, for example, those disclosed by Beattie (WO 95/11755). Any solid surface to which oligonucleotides can be bound, either directly or indirectly, either covalently or non-covalently, can be used. In some

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embodiments, it may be desirable to attach some oligonucleotides covalently and others non-covalently to the same solid support.

A preferred solid support is a high density array or DNA chip. These contain a particular oligonucleotide probe in a predetermined location on the array. Each predetermined location may contain more than one molecule of the probe, but each molecule within the predetermined location has an identical sequence. Such predetermined locations are termed features. There may be, for example, from 2, 10, 100, 1000 to 10,000, 100,000 or 400,000 of such features on a single solid support. The solid support, or the area within which the probes are attached may be on the order of a square centimeter.

Oligonucleotide probe arrays for expression monitoring can be made and used according to any techniques known in the art (see for example, Lockhart et al., Nat Biotechnol 14, 1675-1680 (1996); McGall et al., Proc Nat Acad Sci USA 93, 13555-13460 (1996)). Such probe arrays may contain at least two or more oligonucleotides that are complementary to or hybridize to two or more of the genes described herein. Such arrays my also contain oligonucleotides that are complementary or hybridize to at least 3, 4, 5, 6, 7, 8, 9, 10, 20, 30, 50, 70 or more the genes described herein.

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Methods of forming high density arrays of oligonucleotides with a minimal number of synthetic steps are known. The oligonucleotide analogue array can be synthesized on a solid substrate by a variety of methods, including, but not limited to, light-directed chemical coupling, and mechanically directed coupling (see Pirrung et al., (1992) U.S. Patent No. 5,143, 854; Fodor et al., (1998) U.S. Patent No. 5,800,992; Chee et al., (1998) U.S. Patent No. 5,837,832).

In brief, the light-directed combinatorial synthesis of oligonucleotide arrays on a glass surface proceeds using automated phosphoramidite chemistry and chip masking techniques. In one specific implementation, a glass surface is derivatized with a silane reagent containing a functional group, e.g., a hydroxyl or amine group blocked by a photolabile protecting group. Photolysis through a photolithogaphic mask is used selectively to expose functional groups which are then ready to react with incoming 5° photoprotected nucleoside phosphoramidites. The phosphoramidites react only with those sites which are illuminated (and thus exposed by removal of the photolabile blocking group). Thus, the phosphoramidites only add to those areas selectively exposed from the preceding step. These steps are repeated until the desired array of sequences have been synthesized on the solid surface. Combinatorial synthesis of different oligonucleotide

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analogues at different locations on the array is determined by the pattern of illumination during synthesis and the order of addition of coupling reagents.

In addition to the foregoing, additional methods which can be used to generate an array of oligonucleotides on a single substrate are described in Fodor et al., WO 93/09668. High density nucleic acid arrays can also be fabricated by depositing pre-made or natural nucleic acids in predetermined positions. Synthesized or natural nucleic acids are deposited on specific locations of a substrate by light directed targeting and oligonucleotide directed targeting. Another embodiment uses a dispenser that moves from region to region to deposit nucleic acids in specific spots.

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Hybridization

Nucleic acid hybridization simply involves contacting a probe and target nucleic acid under conditions where the probe and its complementary target can form stable hybrid duplexes through complementary base pairing (see Lockhart et al., WO 99/32660). The nucleic acids that do not form hybrid duplexes are then washed away leaving the hybridized nucleic acids to be detected, typically through detection of an attached detectable label. It is generally recognized that nucleic acids are denatured by increasing the temperature or decreasing the salt concentration of the buffer containing the nucleic acids. Under low stringency conditions (e.g., low temperature and/or high salt) hybrid duplexes (e.g., DNA-DNA, RNA-RNA or RNA-DNA) will form even where the annealed sequences are not perfectly complementary. Thus, specificity of hybridization is reduced at lower stringency. Conversely, at higher stringency (e.g., higher temperature or lower salt) successful hybridization requires fewer mismatches. One of skill in the art will appreciate that hybridization conditions may be selected to provide any degree of stringency. In a preferred embodiment, hybridization is performed at low stringency, in this case in 6x SSPE-T at 37°C (0.005% Triton x-100) to ensure hybridization and then subsequent washes are performed at higher stringency (e.g., 1× SSPE-T at 37°C) to eliminate mismatched hybrid duplexes. Successive washes may be performed at increasingly higher stringency (e.g., down to as low as 0.25× SSPET at 37°C to 50°C) until a desired level of hybridization specificity is obtained. Stringency can also be increased by addition of agents such as formamide. Hybridization specificity may be evaluated by comparison of hybridization to the test probes with hybridization to the various controls that can be present (e.g., expression level control, normalization control, mismatch controls, etc.).

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In general, there is a tradeoff between hybridization specificity (stringency) and signal intensity. Thus, in a preferred embodiment, the wash is performed at the highest stringency that produces consistent results and that provides a signal intensity greater than approximately 10% of the background intensity. Thus, in a preferred embodiment, the hybridized array may be washed at successively higher stringency solutions and read between each wash. Analysis of the data sets thus produced will reveal a wash stringency above which the hybridization pattern is not appreciably altered and which provides adequate signal for the particular oligonucleotide probes of interest.

10 Signal Detection

The hybridized nucleic acids are typically detected by detecting one or more labels attached to the sample nucleic acids. The labels may be incorporated by any of a number of means well known to those of skill in the art (see Lockhart et al., WO 99/32660).

15 Databases

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The present invention includes relational databases containing sequence information, for instance for one or more of the genes of Tables 1-5, as well as gene expression information in various breast tissue samples. Databases may also contain information associated with a given sequence or tissue sample such as descriptive information about the gene associated with the sequence information, descriptive information concerning the clinical status of the tissue sample, or information concerning the patient from which the sample was derived. The database may be designed to include different parts, for instance a sequence database and a gene expression database. Methods for the configuration and construction of such databases are widely available, for instance, see Akerblom et al., (1999) U.S. Patent No. 5,953,727, which is specifically incorporated herein by reference in its entirety.

The databases of the invention may be linked to an outside or external database. In a preferred embodiment, as described in Tables 1-5, the external database is GenBank and the associated databases maintained by the National Center for Biotechnology Information (NCBI).

Any appropriate computer platform may be used to perform the necessary comparisons between sequence information, gene expression information and any other information in the database or provided as an input. For example, a large number of computer workstations are available from a variety of manufacturers, such has those

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available from Silicon Graphics. Client-server environments, database servers and networks are also widely available and appropriate platforms for the databases of the invention.

The databases of the invention may be used to produce, among other things, electronic Northern blots (B-Northerns) to allow the user to determine the cell type or tissue in which a given gene is expressed and to allow determination of the abundance or expression level of a given gene in a particular tissue or cell. The E-northern analysis can be used as a tool to discover tissue specific candidate therapeutic targets that are not overexpressed in tissues such as the liver, kidney, or heart. These tissue types often lead to detrimental side effects once drugs are developed and a first-pass screen to eliminate these targets early in the target discovery and validation process would be beneficial.

The databases of the invention may also be used to present information identifying the expression level in a tissue or cell of a set of genes comprising at least one gene in Tables 1-5 comprising the step of comparing the expression level of at least one gene in Tables 1-5 in the tissue to the level of expression of the gene in the database. Such methods may be used to predict the physiological state of a given tissue by comparing the level of expression of a gene or genes in Tables 1-5 from a sample to the expression levels found in tissue from normal breast tissue, tissue from breast carcinoma or both. Such methods may also be used in the drug or agent screening assays as described herein.

20 Kits

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The invention further includes kits combining, in different combinations, highdensity oligonucleotide arrays, reagents for use with the arrays, signal detection and arrayprocessing instruments, gene expression databases and analysis and database management software described above. The kits may be used, for example, to monitor the progression of breast cancer, to identify genes that show promise as new drug targets and to screen known

and newly designed drugs as discussed above.

The databases packaged with the kits are a typically a compilation of expression patterns from human breast cancer tissue or cell lines and for gene and gene fragments as described herein (corresponding to the genes of Tables 1-5). In particular, the database software and packaged information include the expression results of Tables 1-5 that can be used to predict the cancerous state of a tissue sample by comparing the expression levels of the genes in the tissue or cell sample to the expression levels presented in Tables 1-5.

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The kits may used in the pharmaceutical industry, where the need for early drug testing is strong due to the high costs associated with drug development, but where bioinformatics, in particular gene expression informatics, is still lacking. These kits will reduce the costs, time and risks associated with traditional new drug screening using cell cultures and laboratory animals. The results of large-scale drug screening of pre-grouped patient populations, pharmacogenomics testing, can also be applied to select drugs with greater efficacy and fewer side-effects. The kits may also be used by smaller biotechnology companies and research institutes who do not have the facilities for performing such large-scale testing themselves.

Databases and software designed for use with use with microarrays is discussed in Balaban et al., (2001) U.S. Patent Nos. 6,229,911, a computer-implemented method for managing information, stored as indexed tables, collected from small or large numbers of microarrays, and 6,185,561, a computer-based method with data mining capability for collecting gene expression level data, adding additional attributes and reformatting the data to produce answers to various queries. Chee et al., (1999) U.S. Patent No. 5,974,164, disclose a software-based method for identifying mutations in a nucleic acid sequence based on differences in probe fluorescence intensities between wild type and mutant sequences that hybridize to reference sequences. The object of the method is to predict regions or positions of mutation.

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Without further description, it is believed that one of ordinary skill in the art can, using the preceding description and the following illustrative examples, make and utilize the compounds of the present invention and practice the claimed methods. The preceding working examples therefore, are illustrative only and should not be construed as limiting in any way the scope of the invention.

Examples

Example 1: Preparation of Breast Cancer Profiles

Tissue Sample Acquisition and Preparation

The patient tissue samples were derived from female patients; the average age for the normal and tumor samples was 39 and 52 years respectively. They stem from three different ethnic origins (Caucasian, African-American, and Asian). Furthermore, all tissue samples from Infiltrating Ductal Carcinoa (IDC) patient samples were studied for cancerrelated expression, as 85% of the breast cancer patients were afflicted with this form of the

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disease. The samples are composed of normal, benign, DCIS (ductal carcinoma in-situ), microinvasive, stage L stage II, and stage III breast cancer samples.

Histological analysis of each of the tissue samples was performed and samples were segregated into either normal or malignant categories. The normal tissue samples were acquired from neighboring tissue of patients suffering from one of the following disorders: macromastia, mild fibrosis, infiltrating lobular carcinoma, or infiltrating ducal carcinoma, however; each tissue was diagnosed as normal by histological analysis.

With minor modifications, the sample preparation protocol followed the Affymetrix Gene-Chip Expression Analysis Manual. Frozen tissue was first ground to powder using the Spex Certiprep 6800 Freezer Mill. Total RNA was then extracted using Trizol (Life Technologies). The total RNA yield for each sample (average tissue weight of 300 mg) was 200-500 μ g. Next, mRNA was isolated using the Oligotex mRNA Midi kit (Qiagen). Since the mRNA was eluted in a final volume of 400 μ l, an ethanol precipitation step was required to bring the concentration to 1 μ g/ μ l. Using 1-5 μ g of mRNA, double stranded cDNA was created using the SuperScript Choice system (Gibco-BRL). First strand cDNA synthesis was primed with a T7-(dT₂₄) oligonucleotide. The cDNA was then phenol-chloroform extracted and ethanol precipitated to a final concentration of 1 μ g/ μ l.

From 2 μ g of cDNA, cRNA was synthesized according to standard procedures. To biotin label the cRNA, nucleotides Bio-11-CTP and Bio-16-UTP (Enzo Diagnostics) were added to the reaction. After a 37°C incubation for six hours, the labeled cRNA was cleaned up according to the Rneasy Mini kit protocol (Qiagen). The cRNA was then fragmented (5× fragmentation buffer: 200 mM Tris-Acetate (pH 8.1), 500 mM KOAc, 150 mM MgOAc) for thirty-five minutes at 94°C.

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55 µg of fragmented cRNA was hybridized on the human and the Human Genome U95 set of arrays for twenty-four hours at 60 rpm in a 45°C hybridization oven. The chips were washed and stained with Streptavidin Phycocrythrin (SAPE) (Molecular Probes) in Affymetrix fluidics stations. To amplify staining, SAPE solution was added twice with an anti-streptavidin biotinylated antibody (Vector Laboratories) staining step in between. Hybridization to the probe arrays was detected by fluorometric scanning (Hewlett Packard Gene Array Scanner). Following hybridization and scanning, the microarray images were analyzed for quality control, looking for major chip defects or abnormalities in hybridization signal. After all chips passed QC, the data was analyzed using Affymetrix GeneChip software (v3.0), and Experimental Data Mining Tool (EDMT) software (v1.0).

Gene Expression Analysis

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All samples were prepared as described and hybridized onto the Affymetrix Human Genome U95 array. Each chip contains 16-20 oligonucleotide probe pairs per gene or cDNA clone. These probe pairs include perfectly matched sets and mismatched sets, both of which are necessary for the calculation of the average difference. The average difference is a measure of the intensity difference for each probe pair, calculated by subtracting the intensity of the mismatch from the intensity of the perfect match. This takes into consideration variability in hybridization among probe pairs and other hybridization artifacts that could affect the fluorescence intensities. Using the average difference value that has been calculated, an absolute call for each gene or EST is made.

The absolute call of present, absent or marginal is used to generate a Gene Signature, a tool used to identify those genes that are commonly present or commonly absent in a given sample set, according to the absolute call. For each set of samples, a median average difference was calculated using the average differences of each individual sample within the set. The median average difference typically must be greater than 20 to assure that the expression level is at least two standard deviations above the background noise of the hybridization. For the purposes of this study, only the genes and gene fragments with a median average difference greater than 20 were further studied in detail.

The Gene Signature for one set of samples is compared to the Gene Signature of another set of samples to determine the Gene Signature Differential. This comparison identifies the genes that are consistently present in one set of samples and consistently absent in the second set of samples.

The Gene Signature Curve is a graphic view of the number of genes consistently present in a given set of samples as the sample size increases, taking into account the genes commonly expressed among a particular set of samples, and discounting those genes whose expression is variable among those samples. The curve is also indicative of the number of samples necessary to generate an accurate Gene Signature. As the sample number increases, the number of genes common to the sample set decreases. The curve is generated using the positive Gene Signatures of the samples in question, determined by adding one sample at a time to the Gene Signature, beginning with the sample with the smallest number of present genes and adding samples in ascending order. The curve displays the sample size required for the most consistency and the least amount of expression variability from sample to sample. The point where this curve begins to level off represents the minimum

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number of samples required for the Gene Signature. Graphed on the x-axis is the number of samples in the set, and on the y-axis is the number of genes in the positive Gene Signature.

As a general rule, the acceptable percent of variability in the number of positive genes between two sample sets should be less than 5%.

Fold Change analysis

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The data was first filtered to exclude all genes that showed no expression in any of the samples. The ratio (tumor/normal) was calculated by comparing the mean expression value for each gene in the breast cancer sample set against the mean expression value of that gene in the normal breast sample set. For Table 2, genes were included in the analysis if they had a fold change ≥ 3 in either direction, and a p-value < 0.05 as determined by a two-tail unequal variance t-test. Out of the $\sim 60,000$ genes surveyed by the Human Genome U95 set. 802 genes were present in the overall fold change analysis

15 Expression Profiles of Genes Differentially Expressed in Breast Cancer

Using the above described methods, genes that were predominantly over-expressed in breast cancer, or predominantly under-expressed in breast cancer were identified. Genes with consistent differential expression patterns provide potential targets for broad range diagnostics and therapeutics. For simplicity, applicants examined known genes by hierarchical cluster analysis developed by Eisen and colleagues to determine if functionally related genes would cluster together (see Eisen, et al. Proc Natl Acad Sci USA 95, 14863-14868 (1998)).

Table 2 lists the genes determined to be differentially expressed in cancerous breast tissues compared to normal breast tissue, with the fold change value for each gene. These genes or subsets of these genes comprise an overall breast cancer gene expression profile.

The well-characterized proliferation marker for breast cancer KI-67 had an average-fold change value of 2.8, which was calculated from 15 IDC tissue samples analyzed (see Gerdes, Semin Cancer Biol 1, 199-206 (1990)). As the fold change was below the present 3 fold criteria, the fold change value was not presented in Table 2. Some genes previously shown to be over or under expressed in breast cancer as indicated from the literature such as cytokeratins 5, 14, 15, 17, maspin, MMP 9 and 11, fibronectin, and pituitary tumor transforming 1, etc. are displayed in Table 2 as well as some genes such as p57(kip2), p63/p51/KET, mitosin, and pCDC55 whose expression levels were not previously known to vary in breast cancer.

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The pituitary-tumor transforming 1 gene has been shown to produce in vitro and in vivo tumor-inducing activity (see Zhang et al. Mol Endocrinol 13, 156-66 (1999). In a recent publication, pituitary-tumor transforming 1 has been shown to be over-expressed in mammary adenocarcinomas (see Saez et al. Oncogene 18, 5473-6 (1999)). Also, another study discovered that all 48 colon carcinomas examined over-expressed PTTG1 as compared to normal colorectal tissue, and invasion of the surrounding tissue was associated with higher PTTG1 expression levels (see Heaney et al. Expression of pituitary-tumour transforming gene in colorectal tumours [see comments] Lancet 355, 716-9 (2000)).

Genes listed in Table 2, not reported in the literature to be over-expressed in human breast cancer tissues, include RAD2, FLS353, CKS2, cyclin-selective ubiquitin carrier protein E2-C, ZWINT, Lamin B1 and H2A.X. Although FLS353 has been recently found to be over-expressed in colorectal cancer (see Hufton et al. FEBS Lett 463, 77-82 (1999)), differential expression of FLS353 in breast tumor cells had not been previously demonstrated.

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Cyclin-ubiquitin carrier protein E2-C is another gene over-expressed in breast cancer, which was discovered in this study. Previous studies have shown that when a dominant-negative form of the protein is over-expressed, the mammalian cells arrested in M phase and evidence was provided indicating that this mutant form of cyclin-ubiquitin carrier protein E2-C blocked the destruction of both cyclin A and B (see Townsley et al., Proc Natl Acad Sci U S A 94, 2362-7 (1997)).

The expression levels of the genes in Tables 4 and 5 are associated with various stages of infiltrating ductal carcinoma (Table 4) or infiltrating lobular carcinoma (Table 5). The Tables present the fold change value of expression in the particular disease state compared to normal breast tissue. The genes in these tables may be used alone, or in combination with those listed in Tables 1-3 in the methods, compositions, databases and computer systems of the invention.

Example 2: Diagnostic Subset of Breast Cancer Associated Genes

Table 1 lists the members of a diagnostic subset of genes selected by p-value. This group of genes can be used to differentiate between normal/benign and breast tumor tissue samples including two DCIS samples. Assays using these genes are capable of distinguishing between normal and tumor samples with near 100% efficiency (see Figure 6). Only 1 of the 33 samples shown was misclassified as a normal sample based on the gene

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expression profile when this set of genes was used to analyze the 33 sample set (see Figure 7).

Figures 6 and 7 are three-dimensional plots displaying the relationship of variance derived from gene expression data obtained from patient samples. In Figure 6, normal tissue samples are displayed as darker spheres and the infiltrating ductal carcinoma tissue samples are the lighter spheres. The x-axis represents the first principal component that contains the greatest variance in data of 80%. The y-axis represents the second principal component of 4%. The z-axis represents the third principal component of 3%. Figure 7 displays the results obtained from a separate 33 sample set which is composed of new samples that have no relation to the 28 sample set utilized to discover the gene set of Table 1. Again, the x, y, and z-axes represent the first (63%), second (10%), and third principal components (6%), respectively.

The gene set of Table 1 can thus be used to distinguish normal from cancerous breast tissue.

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Example 3: Myoepithelial and Luminal Cell Marker Genes Examined on a Global Scale

Previous studies have indicated that myoepithelial cells express both epithelial and smooth muscle gene expression markers while luminal epithelial cells fail to express these genes (see Lazard et al., Proc Natl Acad Sci USA 90, 999-1003 (1993)). Cluster analysis identified a group 35 fragments representing 31 genes into one highly correlative cluster and the combination of genes and ESTs are listed in Table 3.

Previous studies have indicated that calponin and myosin heavy chain are expressed in smooth muscle cells and myoepithelial cells while luminal epithelium lack the expression of these genes. Furthermore, the proteins are usually not expressed in invasive ductal carcinoma of the breast (Lazard, et al., supra). Both calponin (fold change—11) and myosin heavy chain (fold change—10.8) were under-expressed in IDC. As indicated in Table 3, other genes associated with smooth muscle that were under-expressed such as smooth muscle gamma-actin, myosin light chain kinase, myosin, heavy polypeptide 11, and Leiomodin 1 and both mysoin polypeptide 11 and leiomodin 1 have not been previously reported to be under-expressed in breast cancer as compared to normal tissue samples.

The expression pattern represented in this particular cluster indicates that a preponderance of tissue samples diagnosed as infiltrating ductal carcinoma exhibit a luminal phenotype while myoepithelial cells were absent. More evidence to support this finding includes the under-expression of cytokeratins 5, 14, 15, and 17 in the tumor samples as

shown in Table 3. Normal myoepithelial cells express cytokeratins 5, 14, 15, and 17 and breast carcinoma cells do not (Trask et al. Proc Natl Acad Sci USA 87, 2319-2323 (1990)). A previous study has indicated that myoepithelial cells are present in normal, benign lesions, grade I infiltrating ductal carcinoma but are absent in carcinomas of grades II and

III (Gusterson et al. Cancer Res 42, 4763-4770 (1982)).

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In addition, components of the basal lamina such as laminin were under-expressed in the infiltrating ductal carcinoma relative to normal tissue samples (Table 3). Both laminin S B3 and laminin-related protein were under-expressed as indicated in Table 3. It has been reported that myoepithelial and basal lamina markers are useful in differentiating microinvasive from ductal carcinomas of the breast (Damiani et al. Virchows Arch 434, 227-234 (1999)).

The set of 35 fragments representing 31 genes as shown in Table 3 could distinguish between intraductal carcinoma and microinvasive DCIS tissue samples based on the disappearance of genes expressed in either basal lamina or myoepithelial cells. There is evidence in the literature that the collapse of the basement membrane as well as the disappearance of an intact myoepithelial cell layer occurs during the invasion process. A multi-gene screen utilizing either of these sets of genes can be used to differentiate between benign and invasive breast neoplasm based on the gene expression fingerprint elucidated in this study.

Figure 8 shows the results of PCA of the 91 sample set with all 35 fragments (representing 31 genes and ESTs) in Table 3. These results demonstrate that PCA using the genes in Table 3 is able to distinguish between non-invasive and invasive breast tissue samples. Figure 8 provides evidence that this group of genes is diagnostically useful for differentiating DCIS samples that are intraductal (non-invasive) from those containing microinvasion. As shown in Figure 8, this group of genes and ESTs is capable of differentiating between two subtypes of DCIS and may constitute a set that is a more sensitive predictor of a microinvasion phenotype.

Example 4: Discovery of Breast Tissue Specific Genes in IDC

Electronic northern (E-northern) analysis determines if a gene of interest is present in a tissue from a database of gene expression information, and if it is present, then at what levels. Expression levels were determined using a GeneChip set that evaluated the expression levels of 60,000 genes in each type of tissue from 28 different normal human tissues. Similar to multi-tissue northern blot analysis, E-northern analysis quickly

determines if a gene of interest is expressed in a particular tissue type and also at what level. E-northern analysis of multiple tissue samples can be evaluated and the determination of exactly how many samples of a particular group that express the gene of interest is tabulated and statistical analysis can be implemented. Multiple samples from the same tissue are not available at this time using conventional multi-tissue northern blot analysis.

The E-northern analysis can be used as a tool to discover tissue specific candidate therapeutic targets that are not over-expressed in tissues such as the liver, kidney, or heart. These tissue types often lead to detrimental side effects once drugs are developed and a first-pass screen to eliminate these targets early in the target discovery and validation process would be beneficial. Furthermore, different tissues have very unique gene expression profiles related to parameters such as proliferation, differentiation, or cell types contained in the tissue that can provide interesting clues into the biological roles of the ESTs.

E-northern analysis was performed for many of the genes clustered in Table 2.

Analysis of the E-northerns revealed that most of the genes were expressed at elevated levels in the thymus. There is high rate of mitosis present in the thymus during T-lymphocyte maturation and many proliferation-associated genes are expressed at elevated levels such as CDC2, cyclin B1, and topoisomerase II alpha. Figure 1 displays the E-northern analysis for topoisomerase II alpha indicating elevated levels of expression in the thymus as compare to the other tissue types detected. Figure 2 shows the results of an E-Northern analysis of transcription factor ICBP90, implicated to be involved with topoisomearse II alpha expression. ICBP90 was also expressed at high levels relative to the other tissue types in the thymus (Figure 2). A study by Hopfner et al. indicated that adult thymus and fetal thymus contained the highest levels of ICBP90 using a 50-tissue RNA dot blot protocol (Hopfner et al. Cancer Res 60, 121-128 (2000)). Most of the genes contained in this cluster contained the highest levels of expression in the thymus.

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Figure 3 shows the results of an E-Northern analysis of the monocarboxylate transporter 4 (MCT4; formerly known as MCT3) which was grouped with genes associated with proliferation. MCT4 is most evident in cells with a high glycolytic rate such as muscle, white blood cells, and tumor cells (Halestrap et al., Biochem J 343 (Pt 2), 281-299 (1999)). A group of multi-tissue northern blots from a recent publication indicate that MCT4 is expressed at high levels in leukocytes but also other tissue types as well (Price et al., Biochem J 329, 321-328 (1998)). Furthermore, electronic-northern analysis indicated high levels of MCT4 were expressed in blood and white blood cells (Figure 3).

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A previously uncharacterized gene only expressed in breast tissue was identified from this study and an E-Northern analysis of the expression pattern of this gene is shown in Figure 4. The distribution pattern of the expression of the gene shows it be used as a marker for breast cancer. The E-northern analysis only displays tissues where the gene of interest is present at detectable levels and breast tissue was the only tissue that this particular gene was under-expressed by -4.2 fold in IDC making it particularly useful as a diagnostic marker.

Another gene that may be used as a diagnostic marker that was not present in a particular cluster is the secreted frizzled-related protein 1. This gene was under-expressed in IDC by -17.7 fold, and the B-northern analysis shown in Figure 5 indicates that it was expressed at greatest levels in breast tissue as well as in the cervix. Using the combination of clustering, fold-change analysis, and B-northern analysis on microarray data one skilled in the art can readily select additional therapeutic and diagnostic markers.

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Although the present invention has been described in detail with reference to examples above, it is understood that various modifications can be made without departing from the spirit of the invention. Accordingly, the invention is limited only by the following claims. All cited patents and publications referred to in this application are herein incorporated by reference in their entirety.

| Top 5 | Table 1: Top 50 by p-value | Genhank | Description | Fold Change | p-values |
|---------|----------------------------|---------------------|--|-------------|-------------------|
| | 37892_at | J04177 | Cluster Incl J04177:Human alpha-1 type XI collagen (COL11A1) mRNA, complete cds (cds=(161,581) // dp=1/9729 Au=Hs,82772 //en=6158 | 10.18941323 | 1.24E-12 |
| 278 35 | 35832_at | AB029000 | Cluster Incl AB029000:Horno sapiens mRNA for KLAA1077 protein, partial cds /cds=(0.2456) /gb=AB029000 /gj=5689490 /ug=Hs.70823 /len=4834 | 4.243716901 | 2.469E-12 |
| 1227 36 | 38566_at | X60382 | Cluster Ind X60382.H.sapiens COL10A1 gene for collagen (alpha-1 type X) /cds=10,2042) /gb=X60382 /gi=30094 /ug=Hs.179729 /len=3198 | 9.164231149 | 4.719E-12 |
| 1226 3 | 38181_at | X57766 | Cluster Incl X57766:Human stromelysin-3 mRNA /cds=(9,1475) /gb=X57766 /gi=456256 /uc=Hs.155324 /len=2247 | 13.96572736 | 5.309E-12 |
| 1159 1 | 1651_at | U73379 | U73379 /FEATURE / DEFINITION=HSU73379 Human cyclin-selective ubiquitin carrier protein mRNA, complete cds | 4.802394237 | 5.8315E-11 |
| 1161 | 33143_s_s | 33143_s_at U81800 | Cluster Incl U61800:Homo sapiens monocarboxylate transporter (MCTS) ImRNA, complete cds /cds=(62,1459), (gb=U8180) (gl=2463633 /(uc+ls,85838 /len=1982 | 0.07793742 | 6.2406E-11 |
| 536 | 34342 <u>_s</u> _s | 34342_s_at AF052124 | Cluster Incl AF052124;Homo sapiens clone 23810 osteopontin mRNA, complete cds /cds=(87,989) /dp=AF052124 /gl=3380431 /lug=Hs,313 /len=1504 | 8.862541971 | 1.1789E-10 |
| 1008 | 31859_at | J05070 | Cluster Incl J05070:Human type IV collagenase mRNA, complete cds /cds=(19,2142)/gb=J05070 /di=177204 /up=Hs. 151738 /len=2334 | 4.66285568 | 1.68 E- 10 |
| 961 | 38116_at | D14657 | Cluster Incl D14657:Human mRNA for KIAA0101 gene, complete cds /cds=(61,396) /gb=D14657 /gl=285938 /ug=Hs.81892 /len=836 | 0.17 | 2.1736E-10 |
| 1007 | 2092_s_at | J04765 | J04765 /FEATURE= /DEFINITION=HUMOSTRO Human osteopontin mRNA, complete cds | 4.119810176 | 3.9853E-10 |
| 277 | 39109_at | AB024704 | Cluster Incl AB024704:Homo sapiens mRNA for fis353, complete cds /cds=(471,2714) /gb=AB024704 /gi=4589928 /ug=Hs, 3329 /len=3403 | 0.228960682 | 6.59E-10 |

| p-values 7.4333E-10 | 7.7763E-10 | 9.3518E-10 | 1.10E-09 | 1.5685E-09 | 1.5906E-09 | 1.7499E-09 | 2.013E-09 | 2.0933E-09 | 2.11E-09 |
|--|--|--|--|--|--|--|---|--|--|
| Fold Change 9.431588747 | 10.43 | 5.975488989 | 6.152409861 | 0.068952382 | 0.139391318 | 0.047728145 | 7.363245733 | 8.54 | 0.233878866 |
| Description Cluster Incl M10905;Human cellular fbronectin mRNA (cds=(0,2383) /gb=M10905 /gl=182696 /uq=Hs,116162 /len=2384 | Cluster Incl Y15915.Homo sapiens mRNA for chimaeric transcript of collagen type 1 ajoha 1 and platelet derived growth factor beta, 1068 bp (css=(0,1067)/gb=Y15915 (gj=2288486 (Lore+)), (lore+), (lore+) | Cluster Incl AA203213:zx57e04.r1 Homo sapiens cDNA, 5 end /done=IMAGE-446622 /done_end=5"/gb=AA203213 /gi=1798923 /ug=Hs.833 /len=879" | Cluster Incl M97938:Human transcription factor ISGF- · 6.152409861 3 mRNA sequence /cds=UNKNOWN /gb=M97936 /ql=475254 /ug=Hs.21486 /len=2607 | Cluster Incl L32137;Human germline oligomeric matrix protein (COMP) mRNA, complete cds Cace=(25,2289) (gb=L32137 /gj=602449 /ug=Hs.1584 /len=2439 | Cluster Incl AL050118:Homo sapiens mRNA; cDNA DKFZp586C201 (from clone DKFZp586C201) fcds=(0,939) (pb=AL050118 (gi=4884143 Nug=1e; 184641 flen=2621 | Cluster Incl AA418080;zv97h07.st Homo sapiens cDNA, 3 end /done=IMAGE-767773 /done_end=3" /gb=AA418080 /gi=2079881 /ug=Hs.3972 /len=543" | Cluster Incl M91670:Human ubiquitin carrier protein (E2-EPF) mRNA, complete cds /cds=(59,738) /dp=M91670 /dp=(1915 /uc=Hs.174070 /len=890 | Cluster Ind AA203476:2x55e01.r1 Homo sapiens cDNA, 5 end /clone=INAAGE-446424 /clone end=5" /gb=AA203476 /gi=1799203 /ug=Hs.159526 /len=828" | Cluster Ind L37747:Homo sapiens lamin B1 gene cds=(340,2100) /gb=L37747 /gj=576839 ug=Hs.89497 len=2849 |
| Genbank M10905 | Y15915 | AA203213 | M97936 | L32137 | AL050118 | AA418080 | M91670 | AA203476 | L37747 |
| <u>Affy</u> 31720_s_at | 35474_s_at Y15915 | 38432_at | 33338_at | 40161_at | 32190_at | 34778_at | 40619_at | 40412_at | 37985_at |
| SeqID 1030 | 1240 | 82 | 1059 | 1021 | 892 | 105 | 1057 | 8 | 1025 |
| # 2 | 55 | 4 | 5 | e , | 11 | 18 | 6 | 20 | 21 |

| p-values 2.35E-09 | 2.42E-09 | 2.6578E-09 | 2.8399E-09 | 3.1699E-09 | 3.56E-09 | 4.0319E-09 | 4.2586E-09 | 5.41E-09 |
|--|--|---|---|--|---|--|--|--|
| Fold Change 7.935291557 | 4.329038319 | 0.233804467 | 0.16 | 0.284181885 | 6.816530863 | 10.36 | 0.103783146 | 4.690939862 |
| Description M13755 /FEATURE=mRNA /DEFINITION=HUMIFN15K Human interferor-induced 17-KDa1/5-KDa protein mRNA, complete cds | Cluster Incl X14850:Human H2A.X mRNA encoding histone H2A.X /cds=(73,504) /gb=X14850 /gj=31972 hin=Hs 147097 /han=1585 | Cluster Ind. X547-921. Asspiens ckshs2 mRNA for Clust protein homologue /cds=(95,334) /gb=X54942 (ni=29978 /trn=Hs 87378 /len=612 | M91670 /FEATURE=/DEFINITION=HUMEZEPI Human ublquitin carder protein (E2-EPF) mRNA, | Custer to K02581:Human thymidine kinase mRNA, complete cds (57,761) /gb=K02581 /gj=339708 /ur=ls 1/f5/07 /len=1421 | Cluster Ind Al375913:tc14c08.x1 Homo sapiens cDNA, 3 end /clone=IMAGE-2063822 /clone_end=3" /gb=Al375913 /gi=4175903 /ug=Hs, 156346 /len=916" | Cluster Ind AF067656:Homo sapiens ZW10 Interactor Zwint mRNA, complete cds /cds=(24,857) /gb=AF067656 /gj=3901271 /ug=Hs.42650 /len=1639 | Cluster Ind AA704137:ag47g01 s1 Homo saplens cDNA, 3 end /clone=IMAGE-1119984 /clone_end=3" /gb=AA704137 /gl=2714055 /ug=Hs.125359 /len=923" | L47276 FFEATURE=UTR#1 IDEFINITION=HUMTOPATR Homo sapiens (cell line HL-50) alpha topolsomerase truncated-form mRNA, 3"UTR |
| Genbank M13755 | X14850 | X54942 | M91670 | K02581 | Al375913 | AF067656 | AA704137 | L47276 |
| Affy 1107_s_at | 40195_at | 40690_at | 893_at | 41400_at | 40145_at | 35995_at | 39395_at | 904_s_at |
| SeqID 1034 | 1221 | 1224 | 1057 | 1010 | 444 | 303 | 205 | 1027 |
| 25 ## | 23 | 24 | 25 | % | | 88 | 59 | 8 |

| p-values 5.4948E-09 | 5.77E-09 | 5.79E-09 | 5.93E-09 | 5.98E-09 | 6.70E-09 | 6.7E-09 | 7.063E-09 | 7.3252E-09 | 8.5667E-09 |
|---|---|--|--|--|--|--|--|--|---|
| Fold Change 0.222082398 6 | 0.0564074 | 0.145966282 | 0.075355998 | 7.624017858 | 7.973012437 | 0.273622601 | 0.139924253 | 12.37 | 0.184984291 |
| Description Cluster Ind AA255922zr85b06.r1 Homo sepiens cDNA, 5 end /done=IMAGE-882451 /done_end=5" /gp=AA255902 /gj=1892406 /ug=Hs.46423 /len=348" | Cluster Incl U05340:Human p55CDC mRNA, complete cds /cds=(110,1609) /gb=U05340 /gi=480031 /ijc=Hs.82906 /lap=1886 | Cluster Incl X02761:Human mRNA for fibronectin (FN pre-cursor) /cds=(0,6987) /gb=X02761 /gj=31396 /ug=Hs, 118162 /len=7680 | Cluster Incl U74612:Human hepatocyte nuclear factor 3ffork head homolog 114 (IFH+114) mRNA complete cds fcds=(114,5519) (gb=U74612 /gj=1842252 Luc+ls.239 len-3474 | Cluster Incl M25753:Human cyclin B mRNA, 3 end /cds=UNKNOWN /gb=M25753 /gj=181243 /ue=Hs. 23860 /len=1452" | Cluster Incl J03464;Human collagen alpha-2 type I mRNA, complete cds, chone pHCOL2A1 cds=(469,4569) (gb=J03464 (gl=179595 Iurels, 179573 (len=5416 | Cluster Incl X15998:H.sapiens mRNA for the chordroitin sulphate proteoglycan versican, V1 splice variant; precursor peptide Acta=(286,7485) /ch=X15998 (pi=37652 /uc=Hs 81800 /len=35224 | Cluster Incl Ac003107:Human DNA from chromosome 19-specific cosmid R30064 containing the COMP gene, genomic sequence Icdse(0,2453) fcb=Ac003107 (di=2623745) lug=18, 1584 (len=2454 | M23263 /FEATURE=/DEFINITION=HUMARB Human androden receptor mRNA, complete cds | Cluster Incl AB020713:Homo sepiens mRNA for KIAA0906 protein, partial cds (cds=(0,2772) /gb=AB020713 /gl=4240300 /ug=Hs.56966 /len=4217 |
| Genbank AA255502 | U05340 | X02761 | U74612 | M25753 | J03464 | X15998 | AC003107 | M23263 | AB020713 |
| Affy 39969_at | 38414_at | 31719_at | 34715_at | 34736_at | 32305_at | 38112 <u>g_</u> at X15998 | 40162_s_at AC003107 | 1577_at | 41812_s_at AB020713 |
| SeqID 91 | 1136 | 1216 | 1160 | 1043 | 1002 | 1222 | 279 | 1041 | 276 |
| # ¥ | 32 | 83 | 8 | 35 | g | 37 | 38 | 33 | 40 |

| Eold Change paralles 30,644 4.165483399 8.99E-09 70,644 9.99E-09 70 70 70 70 70 70 70 70 70 70 70 70 70 | 1.4629E-08 |
|--|---|
| ing 80,644) =907 =907 plete 89 =nrs end=3" 580" TKRA with | 0.17504844 |
| Description Cluster ind US9877-Human low-with CTP-brinding protein (FAB37) mRVA, compete cots forsi-(10,044) gb-u159877 fjel198194 hg-les-220025 harr-907 Cluster ind AF0954481-Homo sapiens putative G protein-coupled receptor (RAIG1) mRVA, compete cots forsi-(10,117) gb-A-1709448 (gi-u169289) Algorithm of the coupled receptor (RAIG1) mRVA, compete compete coupled receptor (RAIG1) mRVA, compete cots, with clustered All repeats in the intronse competered receptor (RAIG1) mroorement and recepto | L BAC CH-1842 149 (BC289730) containing the hERV gene cids=12844,3719 (pg=AC004770 (gi=5212836 hg=Hs.4756 hen=4522 (abc) float/bit saplers in RNA for elengation factor 1 alpha-2 focis=(83,4144) (gi=270940 (gi=58455 hg=Hs.2842 flen=1756 |
| Gentbank U59877 AF095448 AR85552 MI39874 M15205 | X70940 |
| Affy Gents 33371_s_et U59877 33730_st AF9954 1371_s_st M29874 1971_s_st M29874 41583_st AC0047 | 35174 <u> </u> at X70940 |
| 307 307 756 1044 1036 | 1233 |
| ## 4 4 4 4 4 4 4 4 4 4 4 4 4 4 4 4 4 4 | 47 |

| | | | | | • | 10 | | |
|-------------------------------|---------------------|---|---|---|--|---|---|--|
| | b-values | 9.03E-07 | 6.70E-09 | 2.35E-08 | 2.35E-09 | 5.98E-09 | 1.32E-04 | 1.33E-06 |
| | Fold Change | 12.80130327 | 7.973012437 | 7.957396249 | 7.936291557 | 7.624017858 | 7.206250917 | 6.987016934 |
| | Cluster Description | Cluster Incl J04177:Human alpha-1 type XI collager (COL1141) mRNA, complete cds Cds=(161,6581) 405-J04177 (gi=179729 tug=18.82772 /nen=6158 | Ha.179729 Cluster Incl X60382:H.sapiens COL.1041 gene for collagen (alpha-1 type X) (cds=(0,2042) /gb=X60382 /gj=30084 /ug=Hs.179729 /hen=3198 | | U73379 /FEATURE=/DEFINITION=HSU73379 7.936291557 Human cyclin-selective ubiquitin carrier protein mRNA, complete cds | Cluster Incl U81800:Homo sapiens monocarboxylate transporter (Iv(CT3) mRNA, complete cds /cds=(62,1459) (gb=U81800 /gj=2463633 /ug=Hs,85838 Hen=1982 | Cluster Incl. AFD52124: Homo saplens done 23810 osleopomin mRNA, complets cds Icde=(67,689), (gb=AF052124, fgl=3360431 Ag=Hs.313 /len=1504 | Hs.151738 Cluster Incl. U56070:Human type IV collegenase mRVA, complete cds /col=(19,2142) fgb=U6070 fgl=117204 /ug=Hs.151738 len=2334 |
| | Cluster# | Hs.82772 | Hs.179729 | Hs.155324 | Hs.93002 | Hs.85838 | Hs.313 | Hs.151738 |
| | Gene Name | collagen, type XI, alpha 1 | collagen, type X, alpha 1 (Schmid metaphyseal chondrodysplasia) | matrix metalloproteinase 11 (stromelysin 3) | ubiquitin carrier protein E2-C | solute carrier family 16 (monocarboxylic acid transporters), member 3 | secreted phosphoprotein 1 (osteopontin, bone sialoprotein I, early T- lymphocyte activation 1) | matrix metalloproteinase 9 (gelatinase B, 92kD gelatinase, 92kD type IV collagenase) |
| 3-C-D-E | Genbank | J04177 | X60382 | X57766 | U73379 | U81800 | AF052124 | J05070 |
| Table 2: Final Chip A-B-C-D-E | Affy | 37892_at | 38566_at | 38181_at | 1651_at | 33143_s_at | 34342 <u>_s_at8124</u> | 31859_at |
| Table 2: F | SeqID | 1006 | 1227 | 1226 | 1159 | 1161 | 596 | 1008 |
| | #1 | - | 7 | ო | 4 | ιΩ | 9 | ۲ |

| SeqID Affy | | Aff | , | Genbank | Gene Name | Cluster# | Cluster Description | Fold Change | p-values |
|--|-------------------|----------|------------|---|--|-----------|--|-------------|-------------------|
| 1007 2092_s_st J04766 searcreted phosphoprotein to triangular the control of the | 2092_s_at J04765 | J04765 | | secreted phosphoprof (osteopontin sialoprotein lymphocyte: | ein 1 , bone I, early T- activation | Hs.313 | NA786 FFATURE= IDEFINITION=HUMOSTRO Human osteoponfin mRNA, complete cds | 6.188665921 | 1.20E-04 |
| 277 39109_at AB024704 chromosome 20 open reading frame 1 | 39109_at AB024704 | AB024704 | | chromosom reading fran | e 20 open ne 1 | Hs.9329 | Cluster Incl AB024704:Homo sapiens mRNA for Its383, complete cds /cds=(471,2714) /gip=AB024704 /gi=4589928 /ug=Hs.9329 /len=3403 | 6.152409861 | 1.10E-09 |
| 10 1030 31720_s_at M10905 fibronectin 1 | 31720_s_at M10905 | M10905 | M10905 | fibronectin | - | Hs.118162 | Cluster Incl M10906:Human cellular fibronectin mRNA /cds=(0,2383) /gb=M10905 /gj=182696 /ug=Hs.118162 /len=2384 | 6.01925663 | 1.72E-04 |
| 11 1240 35474 <u>, s, at</u> Y15915 collagen, ty | 35474_s_at Y15915 | Y15915 | Y15915 | collagen, ty | pe I, alpha | Hs.172928 | collagen, type I, alpha Hs.172929 Cluster Ind (1985) Scholler Art Stagelers mRNA for dimension and collagen type I alpha and platelet derived growth factor beta, 1088 pp. docs+0.1007) plan*1 (1981 5 gl=2289468 MgHs, 172928 Ment-1068 | 5.903615342 | 1.76E-02 |
| 12 1059 33388_st M97936 signal transducer and activator of transcription 1, 91kD transcription 1, 91kD | 33338_at M97936 | M97936 | | signal transe activator of transcriptior | ducer and | Hs.21486 | Cluster Incl M97836:Human transcription factor ISGF-3 mRNA sequence fods=ÜNKNOWN fgb=M87836 /gi=475254 /ug=Hs.21486 nen-2907 | 5.34214829 | 1.69E-07 |
| 13 1021 40161_at L32137 cardiage oligomenic mathic protein cardiage oligomenic cardiag | 40161_at L32137 c | L32137 C | 0 2 0 10 0 | cartilage oli matrix prote (pseudoach a, epiphyse dysplasia 1 | gomeric iin tondroplasi al auttiple) | Hs.1584 | Cluster Incl 1321371-Human germline oldgomeric mark protein (CoMP) mRNA, complete cds /045268) (gb=32137 /gj=602448 /ug=+1s.1584 /len=2439 | 5.267240726 | 6.09 E- 06 |
| 14 1057 40619_at M91670 ubiquitin carrier protein | 40619_at M91670 | M91670 | | ubiquitin ca protein | mier | Hs.174070 | Hs.174070 Cluster Incl M91670:Human ubiquitin carrier protein (EZ-EPP) mRNA, complete cds (cdc=(59,736) (gb=M91670 (gj=181915 /ug=Hs.174070 /len=390 | 4.694613277 | 2.18E-06 |

| | | | | | 41 | | | | |
|---------------------|--|---|--|---|--|--|---|---|--|
| p-values | 5.41E-09 | 1.68E-10 | 1.97E-06 | 3.67 E- 07 | 1.59E-08 | 9.67E-08 | 8.99E-09 | 0.00077416 | 1.88E-07 |
| Fold Change | 4.690939862 | 4.66285568 | 4.621425831 | 4.486125913 | 4.465375169 | 4.357390421 | 4.165483399 | 4.09 | 4.002408289 |
| Cluster Description | Cluster Incl AA203476:zx55e01.r1 Homo sapiens CDNA, 5 end (Johne=IMAGE-446424 Johne_end=5" /gib=A203476 /gi=1799203 /lug=ls-159626 Aen=828" | Cluster Incl L37747:Homo sapiens lamin B1 gene /cds=(340,2100) /gb=L37747 /gj=576839 /ug=Hs.89497 /len=2849 | M13755 /FEATURE=mRNA IDEFINITION=HUMIFN15K Human interferon- induced 17-402/15-40a protein mRNA, complete cds | Cluster Incl X14850:Human H2AX mRNA encoding histone H2A.X kds=(73,504) fgb=X14850 /gi=31972 /ug=Hs.147097 fenr=1885 | Cluster Incl X54942:H.sapiens ckshs2 mRNA for Cks1 protein hornologue (cds=(65,334) /gb=X54942 /gi=29978 /ug=Hs.83758 /len=612 | Cluster Incl K02581:Human trymidine kinase mRNA, complete cds (cds=(57,761) /dp=K02581 /gj=339708 /ug=Hs.105097 /fer=142.1 | Cluster Ind AF067666:Homo sepiens ZW10 interactor Zwint mRNA, complete cds interact24,857) /gb=AF067656 /gi=3901271 /ug=H6.4260 /len=1639 | M97935 Homo sapiens transcription factor ISGF-3 mRNA, complete cds (MA, MB3 represent transcript regions 5 prime, MiddleA, MiddleA, and 3 prime respectively) | L47276 /FEATURE=UTR#1 //DEFINITION=HUMTOPATR Homo sapiens (cell line HL-60) alpha topoisomerase truncated form mRNA, 3"UTR |
| Cluster# | Hs.252587 | Hs.89497 | Hs.833 | Hs.147097 | Hs.83758 | Hs.105097 | Hs.42650 | Hs.21486 | |
| Gene Name | pituitary tumor- transforming 1 | lamin B1 | interferon-stimulated protein, 15 kDa | H2A histone family, member X | CDC28 protein kinase 2 | thymidine kinase 1, soluble | ZW10 interactor | signal transducer and activator of transcription 1, 91kD | |
| Genbank | AA203476 | L37747 | M13755 | X14850 | X54942 | K02581 | AF067656 | M97935 | L47276 |
| Affy | 40412_at | 37985_at | 1107_s_at | 40195_at | 40690_at | 41400_at | 35995_at | AFFX- HUMISGF3A/ M97935_MA_at | 904_s_at |
| SeqID | 83 | 1025 | 1034 | 1221 | 1224 | 1010 | 303 | 1058 | 1027 |
| #1 | 15 | 9 | 17 | 8 | 19 | 20 | 24 | 52 | ន |

| | p-values | 1.06E-06 | 2.48E-06 | 2.95E-07 | 2.60 E- 08 | 4.11E-03 | 1.28 E -06 | 3.71E-04 |
|---|---------------------|--|--|---|---|---|---|---|
| | Fold Change | 3.870563686 | 3.855167487 | 3.813256493 | 3.80895841 | 3.800908625 | 3.796503387 | 3.781923678 |
| | Cluster Description | Cluster Incl U05340:Human p55CDC mRNA, complete cds /cds=(110,1609) /gb=U05340 /gi=468031 /ug=Hs.82908 /len=1686 | Cluster Incl X02761:Human mRNA for fibronedin (FN precursor) /cds=(0,9987) /dp=X02761 /gj=31396 /ug=Hs.118162 /en=77890 | Cluster Incl JV48/C3-Human headlooyle mucker 3.813259483 leator-3/fork head homolog 11.4 (HFH-11.4) mRNA complete oak Gots-(14.25%) (gb-JV48/12 /ge=1842282 /ug=Hs.23%) Alen=3474 | Cluster incl M25753:Human cyclin B mRNA, 3 end /cds=UNKNOWN /gb=M25753 /gi=181243 /ug=Hs.23960 /len=1452" | Cluster Incl J03464:Human collegen alpha-2 type I mRNA, complete cds, clone pHCOL2A1 (cds=(469,4569) (gb=)03464 (gl=179595 Ng=179573 Inn=5416 | Cluster Incl X15998:H.sapiens mRN4 for the chordrolin sulphate proleoglycan versican, V1 spilce-variant, precursor peptide (cd=(266,7495) gb=X15998 (gj=37662 Clug=Hz, 3180) Men=8224 | Cluster Ind AC003107:Human DNA from cluster Ind AC003107:Human DNA from Cluster Ind S00064 containing the COMP gene, genomic sequence Icase-(0.2453) igb-AC003107 igi-2623749 lug=Hs.1584 /len=2454 |
| ł | Cluster# | Hs.82906 | Hs.118162 | Нs.239 | Hs.23960 | Hs.179573 | Hs.81800 | Hs.1584 |
| | Gene Name | CDC20 (cell division cycle 20, S. cerevisiae, homolog) | fibronectin 1 | forkhead box M1 | cyclin B1 | collagen, type I, alpha Hs.179573 2 | chondroitin sulfate proteoglycan 2 (versican) | cartilage oligomeric matrix protein (pseudoachondroplasi a, epiphyseal dysplasia 1, multiple) |
| | Genbank | U05340 | X02761 | U74612 | M25753 | J03464 | X15998 | AC003107 |
| | Affy | 38414_at | 31719_at | 34715_at | 34736_at | 32305_at | 38112 <u>g</u> at | 40162_s_at AC003107 |
| | SedD | 1136 | 1216 | 1160 | 1043 | 1002 | 1222 | 279 |
| | # | 54 | 22 | 56 | 27 | . 28 | 39 | 90 |

| WO 02/059271 | | 4 | 13 | | PCT/U | IS02/02176 | |
|---|---|--|--|---|---|--|---|
| p-values 9.37E-05 | 1.57E-07 | 1.35E-04 | 5.17E-03 | 1.67E-05 | 9.47E-08 | 3.03E-03 | 1.22E-03 |
| S.74871763 | 3.620640004 | 3.607432569 | 3.556183255 | 3.539960818 | 3.437302377 | 3.435786237 | 3.427509519 |
| Cluster Description M22263 /FEATURE= / DEFINITION=HUMARB Human androgen receptor mRNA, complete ods | AABS1, member RAS Hs.223025 Cluster Incl U59877:Human low-Mr GTP- oncogene family Incling protein (RABS1) mRNA, complete cds loca=(60,544) (gb=U59877 /gj=1388194 log=Hs.223025 flor=907 | Cluster Incl AF095448:Homo sapiens putative G protein-coupled receptor (RAIG1) mRNA, complete cds /cds=(89,1172)/gb=AF095448 /gi=4083889 /ug=Hs.194691 /len=2288 | M29874 /FEATURE= //DEFINITION=HUMCYP2BB Human cytochrome P450-IIB (hIIB1) mRNA, complete cds | M16206 /FEATURE=cds //DEFINITION=HUMMTKRA Human thymidine kinase gene, complete cds, with clustered Atu repeats in the intons | Cluster Ind AC004770:Homo saplens dronnosome 11, BAC OTH-HSP-31168 (BC289730) containing the hFEN1 gene I/ds=(2844,3788) lgb-aC004770 /gj=3212836 /lgr=Hs_47786 llem=452. | Cluster Incl X70940:H.sapiens mRNA for elongation factor 1 alpha-2 /ods=(83,1474) /gb=X70940 /gj=38455 /ug=Hs.2642 /len=1755 | Hs.278613 X67325 /FEATURE=cds /DEFINITION=HSP27 H.sapiens p27 mRNA |
| Cluster# Hs.99915 | Hs.223025 | Hs.194691 | Hs.1360 | Hs.105097 | | Hs.2642 | Hs.278613 |
| Gente Name androgen receptor (dihydrotestosterone receptor, festicular feminization; spinal and bulbar muscular artophy, Kennedy disease) | RAB31, member RAS oncogene family | retinoic acid induced 3 | cytochrome P450, subfamily IIB (phenobarbital- inducible) | thymidine kinase 1, soluble | | eukaryotic translation elongation factor 1 alpha 2 | interferon, alpha- inducible protein 27 |
| Genbank M23263 | U69877 | AF095448 | M29874 | M15205 | AC004770 | X70940 | X67325 |
| Affy 1577_at | 33371_s_at | 33730_at | 1371_s_at | 910_at | 41583_at | 35174 <u>i_</u> at | 425_at |
| <u>SeqID</u> 1041 | 1166 | 307 | 1044 | 1036 | 281 | 1233 | 1231 |
| 3 # # E | 32 | 33 | 8 | 35 | 38 | 37 | 38 |

| | | | | | 44 | | | | |
|---------------------|---|---|--|---|---|---|--|--|---|
| b-values | 4.57E-07 | 1.19E-03 | 0.00448894 | 9.83E-04 | 2.37E-06 | 5.97E-04 | 5.87E-05 | 5.63E-08 | 9.38E-05 |
| Fold Change | 3.42481014 | 3.411920822 | 3.381535863 | 3.370961478 | 3.349219771 | 3.329472506 | 3.328306522 | 3.295678907 | 3.289545724 |
| Cluster Description | Cluster Incl U30872:Human mitosin mRNA, complete cds /cds=(72,9413) /gb=U30872 /g=1000093 /ug=Hs.77204 /len=10189 | Cluster Incl M55153.Human transglutaminase (TGase) mRNA, complete cds (cds=(135,2198) (gb=M55153 /gj=339520 /ug=Hs.8265 /en=3257 | Hs. 180952 X00351 Human mRNA for beta-actin (_5_ M, _3 represent transcript regions 5 prime, Middle, and 3 prime respectively) | Cluster Incl M13509:Human skin collagenase mRNA, complete cds (cds=(68,1477) /dp=M13509 /gi=180664 /ug=H8.83169 /len=1970 | | Cluster Incl X72755:H.sapiens Humig mRNA Icds=(39,416) /gb=X72755 /gi=311375 /ug=Hs.77367 /len=2545 | Cluster Incl D80008:Human mRNA for KIAAN186 gene, complete cds (cds=[94,684) /gb=D80008 /gl=1136431 /ug=Hs.36232 /len=3248 | M25753 /FEATURE=mRNA /DEFINITION=HUMCYCB Human cyclin B mRNA. 3" end | |
| Cluster# | Hs.77204 | Hs.8265 | Hs.180952 | Hs.83169 | Hs.176663 | Hs.77367 | Hs.36232 | Hs.23960 | Hs.184601 |
| Gene Name | centromere protein F (350/400kD, mitosin) | transglutaminase 2 (C polypeptide, protein- glutamine-gamma- glutamyltransferase) | actin, beta | matrix . metalloproteinase 1 (interstitial | For fragment of IgG, low affinity Illa, receptor for (CD16) | monokine induced by gamma interferon | KIAA0186 gene product | cyclin B1 | solute carrier family 7 Hs.184601 (cationic amino acid transporter, y+ system), member 5 |
| Genbank | U30872 | M55153 | X00351 | M13509 | J04162 | X72755 | D80008 | M25753 | M80244 |
| Affy | 37302_at | 38404_at | AFFX- HSAC07/X00 351 M st | 38428_at | 37200_at | 37219_at | 39677_at | 1945_at | 32186_at |
| SedID | 1144 | 1046 | 1215 | 1033 | 1005 | 1234 | 972 | 1043 | 1054 |
| #1 | 93 | 40 | 14 | 42 | 54 | 44 | 45 | 46 | 74 |

| | | | 40 | | | | |
|--|---|--|---|---|---|---|-------------|
| p-values 2.25E-05 | 4.06E-07 | 6.11E-04 | 5.33E-04 | 1.78E-04 | 7.75E-07 | 0.000054 | 0.00000608 |
| Fold Change 3,247818871 | 3.190855222 | 3.147501909 | 3.044487778 | 3.044067136 | 3.036676875 | 0.327422466 | 0.326731583 |
| Cluster Incl M77836:Human pyrroline 5- carboxylate reductase mRNA, complete cds (cds=(11,970) gp=M77836 fgi=189497 Aug+1s:78277 flem=1782 | Cluster Incl M63199:Human platelet-derived endothelial cell growth factor mRNA, complete case fodes (123.1671), pgb=M63193 /gl=189700 hin=1-8, 73948 /hen=1-587 | Cluster Incl U70370; Human hindlimb expressed homeobox protein backfoot (Bft) mRNA, complete cds (cas=(111,1055) fgb=U70370 fgl=1870670 /ug=Hs.84136 | Hs.105440 Cluster Incl U39940:Human hepatocyte nuclear factor-3 alpha (HNR-3 alpha) mRNA, complete cus /cds=(87,1509) /gb=U39840 /gl=1086121 /nm=48-115440 /me=2872 | Cluster Incl L29254:Human (clone P1-5) L- iditol-2 dehydrogenase gene /cds=(137,1210) /gb=1.29254 /gj=808013 /ug=Hs.878 /len=2519 | X05380 /FEATURE=cds /DEFINITION=HSCDC2 Human CDC2 gene involved in cell cycle control | 0 10 10 - | |
| Cluster# Hs.79217 | Hs.73946 | Hs.84136 | Hs.105440 | Hs.878 | Hs.184572 | Hs.135281 | Hs.22174 |
| Gene Name pyrroline-5- carboxylate reductase | endothelial cell growth factor 1 (platelet-derived) | paired-like homeodomain transcription factor 1 | hepatocyte nuclear factor 3, alpha | sorbitol dehydrogenase | cell division cycle 2, G1 to S and G2 to M | aipha-actinin-2- associated LIM protein | |
| Genbank M77836 | M63193 | U70370 | U39840 | L29254 | X05360 | AF002282 | U50527 |
| Affy 37741_at | 36879_at | 37920_at | 37141_at | 38763_at | 1803_at | 39690_at | 1527_s_at |
| SeqID 1053 | 1049 | 1158 | 1149 | 1020 | 1219 | 285 | 1153 |
| ## 8 | 49 | 92 | 51 | 25 | 53 | 54 | 55 |

| | | | | | | 40 | | | | |
|---|---------------------|---|---------------------------------------|--|---|--|---|--|--|--|
| | p-values | 0.0000229 | 0.000191 | | 0.0001 | 2.84E-08 | 0.000642 | 0.00000552 | 0.0000133 | 0.000834 |
| | Fold Change | 0.326719388 | 0.326431324 | | 0.323566748 | 0.323306977 | 0.323187581 | 0.323178779 | 0.321819387 | 0.3197207 |
| } | Cluster Description | Cluster Incl AF001691:Homo sapiens 195 kDa conflide envelope precursor mRNA, complete cost scds=(90,5360) (gb=AF001691 fgl=2188845 (ug=Hs,74304 flen=8227 | U84487 /FEATURE= /DEFINITION=HSU84487 | Human CX3C chemokine precursor, mRNA, alternatively spliced, complete cds | Cluster Incl D84110:Homo sapiens mRNA for RBP-MS/lype 4, complete cds /cds=(566,1156) /gb=D84110 /gi=1699552 /ug=Hs.80248 /len=1594 | Cluster Incl D14686:Human gene for glyche cleavage system T-protein (cds=(146,1356) /gb=D14686 (gi=994760 /ug=Hs.102 /len=2119 | Cluster Incl J02876:Human placental folate binding protein mRNA, complete cds /cds=(262, 1029) /db=J02876 /gl=182413 /ug=Hs.24194 /fen=1211 | S82539 /FEATURE= /DEFINITION=S82539 insulin receptor substrate-1 [human, skeletal muscle, mRNA, 5828 nt] | Cluster Incl X67098:H.sapiens rTS alpha mRNA containing four open reading frames icds=UNIKNOWN /gb=x67098 /gi=475908 /ug=Hs.180433 fan==1817 | Cluster Incl U58516:Human breast epithelial antigen BA46 mRNA, complete cds /cds=(60,1223)/gb=U58516 /gi=1381161 /ug=Hs.3745 /len=1934 |
| ! | Cluster # | Hs.74304 | Hs.80420 | | Hs.80248 | Hs.102 | Hs.24194 | Hs.96063 | Hs.180433 | Hs.3745 |
| | Gene Name | periplakin | small inducible | cytokine subfamily D (Cys-X3-Cys), member 1 (fractalkine, neurotactin) | RNA-binding protein gene with multiple splicing | aminomethyltransfera se (glycine cleavage system protein T) | folate receptor 2 (fetal) | insulin receptor substrate 1 | rTS beta protein | milk fat globule-EGF factor 8 protein |
| | Genbank | AF001691 | U84487 | | D84110 | D14686 | J02876 | S62539 | X67098 | U58516 |
| | Affy | 36890_at | 823_at | | 38049 <u>g</u> at | 41120_at | 33871_s_at | 851_s_at | 33263_at | 34403_at |
| | SegID | 284 | 1164 | | 973 | 963 | 1001 | 1120 | 1230 | 2 |
| | #1 | 28 | 22 | | 28 | 20 | 09 | 61 | 62 | 93 |

| | | | 47 | | | |
|---|--|--|--|---|---|---|
| 0.000387 | 0.00529 | 0.00523 | 0.000012 | 0.0224 | 0.0000881 | 0.000764 |
| Fold Change 0.319370936 | 0.317801655 | 0.314829644 | 0.313476219 | 0,313295747 | 0.312357015 | 0.31152273 |
| Cluster Instantion Cluster Description Cluster Inst V00588:Human mRN4 encoding the c-myc oncogene fods=(568,1877) (gb=V00568 /gj=34815 /kg=Hs.79070 | Cluster Incl Mz2234.Human amhopeptidase VIVCD13 m444 encoding aminopeptidase N, complete ods cfas+(120,3023) (gp=HZ2324 /gj=178835 /ug=Hs,1239 /ten=3477 | prominin (mouse)-like Hs.112360 Cluster Incl AF027208:Homo sapiens AC133 artigen artigen mittyen mittyen met artigen selected (ASS4) (492–4607208 (492–46.11236) Included Incl | D00632 /FEATURE= //DEFINITION=HUMGSHPXA Homo sapiens mRNA for glutathione peroxidase, complete cds | L22624 /FEATURE=expanded_cds /DEFINITION=HUMIMATRY06 Human matrilysin gene, exon 6 and complete cds | Cluster Ind Z26653:H.sapiens mRNA for lamlinh M chain (merosih) /cds=(49,9381) /gb=Z2663 /gi=438055 /ug=Hs.75279 /fen=8534 | L13740 /FEATURE= /DEFINITION=HUMTR3A Human TR3 orphan receptor mRNA, complete cds |
| Cluster# Hs.79070 | Hs.1239 | Hs.112360 | Hs.172153 | Hs.2256 | Hs.75279 | Hs.1119 |
| Gene Name v-myc avian myelocyfomatosis viral oncogene homolog | alanyl (membrane) aminopeptidase (aminopeptidase N, aminopeptidase M, microsomal aminopeptidase, CD13, p150) | prominin (mouse)-like | glutathione peroxidase 3 (plasma) | matrix metalloproteinase 7 (matrilysin, uterine) | laminin, alpha 2 (merosin, congenital muscular dystrophy) | nuclear receptor subfamily 4, group A, member 1 |
| Genbank V00568 | M22324 | AF027208 | D00632 | L22524 | Z26653 | L13740 |
| Affy. 37724_at | 39385_at | 41470_at | 770_at | 668_s_at | 36917_at | 279_at |
| SeqID 1169 | 1040 | 292 | 828 | 1018 | 1244 | 1016 |
| # 99 | 99 | 99 | 67 | 88 | 69 | 02 |
| | | | | | | |

| | | | 40 | | | | |
|--|--|---|--|--|--|-------------------|------------------|
| p-values 0.0000193 | 0.00000415 | 0.000274 | 0.000239 | 0.0000802 | 1.85E-07 | 7.42E-08 | 0.00149 |
| 6.310894189 | 0.308897008 | 0.306961223 | 0.303827048 | 0.303503697 | 0.300299695 | 0.297519832 | 0.296769339 |
| Cluster Incl AFORZ868: Home seplent gamma Cluster incl AFORZ868: Home seplent gamma butynotherine hydroxylase (GBH) mRNA, complete cds (cds=(68,1229) (gb=AF08288) (gl=3746904 hg=Hs.9697 her=1584 | Cluster Incl AF016004:Homo sapiens m6b1 mRNA, complete cds /cds=(254,1051) /gb=AF016004 /gj=3387766 /ug=Hs.78361 | | U04313 /FEATURE= /DEFINITION=HSU04313 0.303827048 Human maspin mRNA, complete cds | Cluster Ind Z19574:H.sapiens gene for cytokeratin 17 /cds=(64,1362) /gb=Z19574 /ni=30378 /in=Hs_2785 /len=1518 | | | |
| Cluster# Hs.9667 | Hs.5422 | Hs.285857 | Hs.55279 | Hs.2785 | Hs.106070 | Hs.64016 | Hs.155376 |
| Gene Name butyrobetaine (gamma), 2- oxoglutarate dloxygenase (gamma- butyrobetaine hydroxylase) | glycoprotein M6B | franscription factor 7- like 2 (T-cell specific, HMG-box) | protease inhibitor 5 (maspin) | keratin 17 | cyclin-dependent kinase inhibitor 1C (p57, Kip2) | protein S (alpha) | hemoglobin, beta |
| Genbank AF082868 | AF016004 | Y11306 | U04313 | Z19574 | U22398 | M15036 | L48215 |
| Affy 38339_at | 37251_s_at AF016004 | 32025_at | 863_g_at | 34301_r_at | 39545_at | 35752_s_at M15036 | 32052_at |
| SeqID 305 | 290 | 1239 | 1136 | 1242 | 1143 | 1035 | 1028 |
| # 7 | 22 | ಜ | 74 | 75 | 76 | 4 | 78 |

| | | 4 | 9 | | | |
|---|---|---|---|---|---|---|
| 0.0000556 | 0.00000184 | 0.00999 | 0.0000096 | 0.00688 | 8.07E-07 | 0.000483 |
| Fold Change 0.295641772 | 0.29509428 | 0.29444252 | 0.292877726 | 0.29048531 | 0.2896161 | 0.288749983 |
| Cluster Description Cluster Ind AF022383-thon septems LIM domain briding protein (LDB1) mRNA, complete cds (cds-(254,1297) /gb=AF052389 /gj=3044085 /ug=Hs.4890 /len=2398 | Hs.169172 Cluster Incl AJ000388:Homo sapiens mRNA for calpain-like professe CANPX /cds=(182,2107) /gb=AJ000388 /gi=2274961 /uc=Hs.169172 /len=3617 | Cluster Incl M12272-Horno sepiens alcohol cluster Incl M1227-Horno sepiens alcohol ebbydrogenase class I gamma subunit (ADH3) mRNA, complete cab (xds=(60,1207) (gb=M12272, gj=178147 lug=Hs.2523 /len=1486 | Hs. 184222. Cluster Into Usb2G77-Homo saplens down syndome candidate region 1 (DSCR1) gard, afternative exon 1, complete cals rota=(8,457). (ga=08,227) (ga=08,1287 / ug=41s,184222.) flen=227. | Cluster Incl AF098462:Homo saplens stanniocalcin-related protein mRNA, complete cas fous=(134,1042),/gb=AF098462/g=4050037 /ug=Hs.155223./en=2380 | Cluster Incl AL021977-thK447C4.1 (novel MAFF (v-maf musculoaponaurotic inbrosaroma (avelani) onogane family, protein F) LIKE protein) Cdc=(0,494) /gb=AL021977 /gj=4944526 /ug=Hs,51305 /arr=2128 | Cluster Incl Z22865.H.sapiers dermatopontin mRNA, complete CDS /cds=(12,617) /gb=Z22865 /gj=311613 /ug=Hs.80552 /len=729 |
| Cluster # Hs.4980 | Hs.169172 | Hs.2523 | Hs.184222 | Hs.155223 | Hs.51305 | Hs.80552 |
| Gene Name LIM domain binding 2,polymyositis/sclerod erma autoantigen 2 (100kD) | calpain-like protease | alcohol dehydrogenase 1 (class I), alpha polypeptide,alcohol dehydrogenase 3 (class I), garmna | Down syndrome oritical region gene 1 | stanniocalcin 2 | v-maf musculoaponeurotic fibrosarcoma (avian) oncogene family, protein F | dermatopontin |
| Genbank AF052389 | AJ000388 | M12272 | U85267 | AF098462 | AL021977 | Z22865 |
| Affy 36065_at | 40475_at | 36247_f_at | 32168_s_at | 32043_at | 36711_at | 38059_g_at |
| SeqID 298 | 848 | 1031 | 1165 | 308 | 851 | 1243 |
| ## 62 | 80 | 26 | 82 | 88 | 84 | 82 |

| WO 02/05 | 9271 | | | 50 | | | PCT/US | 02/02176 |
|---|---|------------------------------------|-----------------------------------|--|--|---|---|--|
| p-values 0.00000058 | 0.000216 | 0.0000355 | 0.00679 | 0.000174 | 0.0000231 | 0.000495 | 0.00000039 | 0.000161 |
| Fold Change 0.288106064 | 0.287855932 | 0.287831724 | 0.287696584 | 0.285399252 | 0.283121329 | 0.282677481 | 0.281207961 | 0.279338963 |
| Cluster Incl. 104111:Human c-jun proto oncogene (July Complete cds, clone hCL1 (date=[974,1969) /gb=.04111 /gj=196624 | L24203 FEATURE - DEFINITION=HUMDK Homo sapiens ataxia-telangiectasia group D- associated ordelin mRNA, comolete cds | | | L19871 /FEATURE= //DEFINITION=HUMATF3X Human activating transcription factor 3 (ATF3) mRNA, complete cds | Cluster Incl U31384:Human G protein gamma- 11 suburul mRNA, complete cds /cds=(107,328) /gb=U31384 /gi=995920 /to=H5,83381 /len=622 | Cluster Incl AB002409:Homo sapiens mRNA for SLC, complete cds /cds=(56,462) /gb=AB002409 /gl=2335034 /ug=Hs,57907 /fen=852 | | Hs.211682 Clustor Incl U489569-Homo sapiens myosin light 0.279339963 chain fund myosin light 0.279339963 chain fund myosin light 0.279339963 Acd=119.248959 (gi=1377819 /ug=1s.211592 Ann=5926 |
| Cluster # | Hs.82237 | Hs.173859 | Hs.171695 | Hs.460 | Hs.83381 | Hs.57907 | Hs.32309 | Hs.211582 |
| Gene Name v-jun avian sarcoma virus 17 oncogene homolog | ataxia-telangiectasia group D-associated | frizzled (Drosophila) homolog 7 | dual specificity phosphatase 1 | activating transcription factor 3 | guanine nucleotide binding protein 11 | small inducible cytokine subfamily A (Cys-Cys), member 21 | inositol polyphosphate-1- phosphatase | myosin, light polypeptide kinase |
| Genbank J04111 | 124203 | AB017365 | X68277 | L19871 | U31384 | AB002409 | L08488 | U48959 |
| Affy 32583_at | 1898_at | 33222_at | 1005_at | 287_at | 37908_at | 36503_at | 656_at | 32847_at |
| SeqID 1004 | 1019 | 274 | 1232 | 1017 | 1145 | 270 | 1013 | 1151 |

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|---------------------|---|---|---|---|--|--|--|--|
| b-values | 0.00000917 | 0.00000688 | 0.000014 | 6.7E-09 | 0.0000349 | 4.33 E -07 | 0.00000159 | 0.000228 |
| Fold Change | 0.278228021 | 0.276610822 | 0.273963608 | 0.273622601 | 0.26818697 | 0.267616256 | 0.263585259 | 0.260058306 |
| Cluster Description | Cluster Incl S62539:insulin receptor substrate- 1 (humen, skeletal muscle, mRN4, 5828 nt] /cds=(1020,4748) (pt=562539 /gi=386256 /ug=Hs,96063 /en=5799 | U22398 /FEATURE= /DEFINITION=HSU22398 Human Cdk-Inhibitor p57KIP2 (KIP2) mRNA, complete cds | Cluster Incl U07919:Human aldehyde dehydrogenase 6 mRNA, complete cds /cds=(52,1590)/gb=U07919/gj=995897 /ug=Hs.75746/len=3442 | Cluster Incl U35139:Human NECDIN related protein mRNA, complete ods /cds=(58,1023) /gb=U35139 /gl=1754970 /ug=Hs.50130 /en=1592 | Cluster Incl M68840:Human monoamine oxidase A (MACA) mRNA, complete cds Cds=(73,1656) /gb=M6840 /gi=187352 /lg=fs:183109 /len=1949 | Cluster Incl AF093118:Homo saplens UP50 mRNA, complete cds (cds=(188,1514) (gb=AF093118 /gi=3676821 /ug=Hs.11494 flen=2019 | Cluster Incl U10492:Human Mox1 protein (MOX1) mRNA, complete cds (cds=(29,793) /gb=U10492 /g=505653 /ug=Hs.438 /len=2315 | Cluster Ind AF022797.Homo sapiens Intermedite conductance calcium-activated potassium channel (INCAs) mRNA, complete cds (cds=(396,1679) /gb=AF022797 /gj=267.4356 /ug=Hs. 10082 /len=2238 |
| Cluster# | Hs.96063 | Hs.106070 | Hs.75746 | Hs.50130 | Hs.183109 | Hs.11494 | Hs.438 | Hs.10082 |
| Gene Name | insulin receptor substrate 1 | cyclin-dependent kinase inhibitor 1C (p57, Kip2) | aldehyde dehydrogenase 6 | necdin (mouse) homolog | monoamine oxidase A | fibulin 5 | mesenchyme homeo box 1 | potassium intermediate/small conductance calcium- activated channel, subfamily N, member |
| Genbank | S62539 | U22398 | U07919 | U35139 | M68840 | AF093118 | U10492 | AF022797 |
| Affy | 41049_at | 1787_at | 36686_at | 36073_at | 41772_at | 39038_at | 36010_at | 41106_at |
| SeqID | 1120 | 1143 | 1138 | 1147 | 1050 | 306 | 1139 | 291 |
| ## | 92 | 96 | 26 | 86 | 66 | 100 | 101 | 102 |

| | | | | | 52 | | | |
|---------------------|--|--|--|--|--|---|--|--|
| p-values | 4.12E-07 | 0.0000904 | 0.0000232 | 0.0025 | 0.00000113 | 0.000041 | 0.00041 | 0.00105 |
| Fold Change | 0.260055335 | 0.259339901 | 0.259264106 | 0.258780062 | 0.258230155 | 0.257792937 | 0.256900234 | 0.256815942 |
| Cluster Description | Cluster Ind U17760:Human laminin S B3 chain (LAMB3) gene /cds=(389,3917) /gj=U17760 /gj=2182192 /ug=Hs.75517 /len=4213 | M98539 /FEATURE=exon /DEFINITION=HUMPDS03 Human prostaglandin D2 synthase gene, exon 7 | Cluster Incl W61005:zd29a11.s1 Homo sapiens CDNA, 3 and /olone=INAGE:342044 /clone_end=3" /gb=W6100,5 /gi=1367764 /lug=ls,14896 /fen=786" | Cluster Incl J00153:Human alpha globin gene cluster on chromosome 16- zeta gene fcd==(0,428) /gb==100153 /gi==183794 /ug=He, 182374 /len=429 | Cluster Ind U61374:Human novel protein with short consensus repeats of 8ix cystelines mRNA, complete cds /cds=(41,1426) /gb=U61374 /gj=1778409 /ug=Hs.15154 Inn=1800 | Cluster incl X57025:Human IGF-I mRNA for insulin-like growth factor I /ods=(186,627) /gj=X57025 /gj=33007 /ug=Hs.85112 /en=7238 | Hs.180878 Cluster Incl M15856:Human lipoprotein lipase mRNA, complete cds /cds=UNIXNOWN /gb=M15856 /gj=187209 /ug=Hs.180878 /fen=3849 | Cluster Ind X00129:Human mRNA for retinol binding protein (RBP) (xds=f\$1,580) /gb=X00129 /gi=35896 /ug=Hs. 76461 /len=882 |
| Cluster# | Hs.75517 | Hs.8272 | Hs.14896 | Hs.251577 | Hs.15154 | Hs.85112 | Hs.180878 | Hs.76461 |
| Gene Name | laminin, beta 3 (nicein (125kD), kalinin (140kD), BM600 (125kD)) | prostaglandin D2 synthase (21kD, brain) | DHHC1 protein | hemoglobin, alpha 1 | sushi-repeat- containing protein, X chromosome | insulin-like growth factor 1 (somatomedia C) | lipoprotein lipase | retinol-binding protein 4, interstitial |
| Genbank | U17760 | M98539 | W61005 | J00153 | U61374 | X57025 | M15856 | X00129 |
| Affy | 36929_at | 216_at | 39750_at | 31525_s_at | 31855_at | 38737_at | 41209_at | 32552_at |
| SealD | 1141 | 1060 | 1191 | 666 | 1156 | 1225 | 1037 | 1214 |
| ## | 103 | 401 | 105 | 106 | 107 | 108 | 109 | 110 |

| | | | | | | | | _ | |
|---|---------------------|---|---|--|---|---|--|---|--|
| | p-values | 0.000837 | 0.0321 | 0.000106 | 0.000998 | 0.000037 | 0.0000523 | 0.0000179 | 0.000763 |
| | Fold Change | 0.256739005 | 0.251627664 | 0.251471671 | 0.251434843 | 0.250173978 | 0.249382262 | 0.248516754 | 0.242494771 |
| | Cluster Description | Cluster Incl U41518:Human channel-like integral membrane protein (ACP-1) mRNA, clone ACP-1-2344, partial cds (cds=(0,460) (dp=U41518 (gi=1314305 /ug=Hs.74502 Inen=2344 | Cluster Ind Y10179:H.sapiens mRNA for protectir-inducible protein (cds={36,476}) (gb=Y10179 (gi=2282895 /ug=Hs.98949 flen=576 | Cluster Incl K03000:Human aldehyde dehydrogenase 1 mRNA kds=(0,1022) kgb=(03000 kgl=178399 kug=Hs.76392 flen=1560 | Cluster ind M25079:Human sickle cell betaglobin mRNA, complete cds /ds=(0,443) /dp=M25079 /gi=179408 /ug=Hs.234764 | Cluster Incl U39447:Human placenta copper monamine oxidase mRNA, complete cds icds=(160,2451) igb=U39447 ig=1399031 iug=Hs, 198241 ilen=4023 | | L07594 /FEATURE= //DEFINITION=HUMTGFB3C Human transforming growth factor-bets type III receptor (TGF-beta) mRNA, complete cds | Cluster Ind U05981; Human hepatic dihydrodiol 0.242494771 dehydroganese gene fots=(26,987) geb—105881 (gi=487134 Mg=He.201967 An=1022 |
| į | Cluster# | Hs.74602 | Hs.99949 | Hs.76392 | Hs.155376 | Hs.198241 | Hs.85112 | Hs.79059 | Hs.275374 |
| | Gene Name | aquaporin 1 (channel- forming integral protein, 28kD) | prolactin-induced protein | aldehyde dehydrogenase 1, soluble | hemoglobin, beta | amine oxidase, copper containing 3 (vascular adhesion protein 1) | insulin-like growth factor 1 (somatomedia C) | transforming growth factor, beta receptor III (betaglycan, 300kD) | aldo-keto reductase family 1, member C1 (dilhydrodisol dehydrogenase 1; 20- aipha (3-alpha)- hydroxysteroid dehydrogenase) |
| | Genbank | U41518 | Y10179 | K03000 | M25079 | U39447 | X57025 | L07594 | U05861 |
| | Affy | 36156_at | 41094_at | 37015_at | 31687_f_at | 33756_at | 1501_at | 1897_at | 32805_at |
| | SedID | 1150 | 1238 | 1011 | 1042 | 1148 | 1225 | 1012 | 1137 |
| | # | 1 | 112 | 113 | 1 | 115 | 116 | 117 | 11 |

| 77 0 021007271 | | 54 | | | 101/0502021/0 | | |
|--|--|---|---|--|--|--|-------------|
| <u>p-values</u> 2.11E-09 0.0000034 | 3.37E-08 | 6.59E-10 | 0.000415 | 0.031 | 3.52E-07 | 0.00000788 | 0.0000975 |
| Fold Change 0.233878866 0.233549169 | 0.231782987 | 0.228960682 | 0.228917694 | 0.226749627 | 0.225166891 | 0.222238905 | 0.220336613 |
| Cluster Description Cluster Ind U81992: Hone spelles C2H2 zinc finger protein PLAG1. (PLAG1.1) mRNA, complete cds (cds=(176,141),1gb=U81992 glj=5513-452,020,pells.7528,561er=2681 Cluster Ind X9351 G01, sapilers mRNA for 37 kDe LIM domain protein (cds=(41,1027) glp=X83610 (gl=1085021 /ug=1s.78981 filen=130 | Cluster Incl AF039843:Homo sapiens Sprouty 2 (SPRY) mRN4, complete cds (cd=(390,1337) (4b=AF039843 /gl=2809399 /tucHs, 18676 (hen=2117 | Cluster Incl L10373:Human (done CCG-B7) mRNA sequence fcds=UNKNOWN fgb=L10373 fgi=307287 fug=Hs.82749 flen=1792 | D10367 /FEATURE= DEFINITON=HUMMHCAAA Homo saplens mRNA for smooth muscle myosin heavy chain, partial cds. | Cluster Ind U33147:Humen mammeglobin mRNA, complete cds /cds=(60,341) /gb=U33147 /gl=1199595 /ug=Hs.46452 /en=503 | Cluster Incl. J04076:Human early growth response 2 protein (EGR2) mRNA, complete cds fods=(203,1423), gb=J04076 /gj=181986 /tuc=ls, 1365 /len=2700 | | |
| Cluster # Hs.75825 Hs.79691 | Hs.18676 | Hs.82749 | | Hs.46452 | Hs.1395 | Hs.137569 | Hs.119651 |
| Gene Name pleiomorphic adenoma gene-like 1 LIM domain protein | sprouty (Drosophila) homolog 2 | transmembrane 4 superfamily member 2 | | mammaglobin 1 | early growth response 2 (Krox-20 (Drosophila) homolog) | tumor protein 63 kDa with strong homology to p53 | glypican 3 |
| <u>Genbank</u> U81992 X93510 | AF039843 | L10373 | D10667 | U33147 | J04076 | Y16961 | U50410 |
| Affy 36943_r_at 32610_at | 33700_at | 38408_at | 774_g_at | 36329_at | 37863_at | 31791_at | 39350_at |
| 1162 1137 | 294 | 1014 | 096 | 1146 | 1003 | 1241 | 1162 |
| # 1 150 | 121 | 122 | 123 | 124 | 125 | 126 | 127 |

| WO 02/03927 | 1 | | 55 | | PC1/US02/021/0 | | |
|--|---|---|--|---|--|--|--|
| p-values 0.00000138 | 9.64E-09 | 0.0000527 | 0.0000209 | 0,00000443 | 0.000166 | 0.0000371 | 4.51E-07 |
| Fold Change 0.218509986 | 0.216723881 | 0.216083178 | 0.207358276 | 0,206077576 | 0.202234909 | 0,19705452 | 0.196869236 |
| Cluster IncXA4T22-Human mRNA for e 64 Kd automitigen expressed in thyroid and extra- coular musche (date/12,1360) (gbxX6+162 (gl=28968 /ug=Hs, 78386 /lorr=3849 | Cluster Incl U69263;Human matrilin-2 precursor mRNA, partial cds /cds=(0,941) /gb=U69263 /gi=2072789 /ug=Hs.19368 | | Cluster Incl M38820:Human cytokine (GRO-bet) mRNA, complete cds (Cds=(74,397) fgb=M38820 /gi=183628 /ug=Hs.75765 flen=1110 | Cluster Incl U19495:Human intercrine-alpha (InIRI) mRNA, complete cds (rds=(473,742) /gb=U19495 /gi=1754834 /ug=Hs.169672 | | | Cluster ind L13463:Human helix-loop-helix basic phosphoprotein (G0S8) mRNA, complete cds /cds=(32,667) /gb=L13463 /gi=292054 /ug=Hs.78944 /len=1345 |
| Cluster# Hs.79386 | Hs.19368 | Hs.118223 | Hs.75765 | Hs.237356 | Hs.251754 | Hs.110903 | Hs.78944 |
| Gene Name leiomodin 1 (smooth muscle) | matrilin 2 | microfibrillar- associated protein 4 | GRO2 oncogene | stromal cell-derived factor 1 | secretory leukocyte protease inhibitor (antileukoproteinase) | claudin 5 (transmembrane protein defeted in velocardiofacial syndrome) | regulator of G-protein signalling 2, 24kD |
| <u>Genbank</u> X54162 | U69263 | L38486 | M36820 | U19495 | X04470 | AF000959 | L13463 |
| Affy 37765_at | 32239_at | 39066_at | 37187_at | 32666_at | 32275_at | 38995_at | 37701_at |
| SeqID 1223 | 1157 | 1026 | 1045 | 1142 | 1218 | 782 | 1015 |
| 128 | 129 | 130 | 13 | 132 | 133 | 134 | 135 |

| p-values | 0.0108 | 2.64E-10 | 0.000818 | 4.71E-08 | 0.00058 | 9.21E-08 |
|---------------------|--|--|---|--|---|---|
| Fold Change | 0.195015273 | 0.191432122 | 0.19139473 | 0.187743879 | 0.187570884 | 0.186647173 |
| Cluster Description | Cluster Ind LUGRECH-forms sapiers car2b (car2), car2e (car2e (car | Cluster ind M18653:+Omo saplens dystrophin (DMD) mRNA, complete das fods=UNKACOWN (gb-M18653 fg=18186 fug=Hs.169470 Aen=13857 | Cluster Incl AF044311:Homo sepiens gamma- synuciein gene, complete cds /cds=(52,435) /gb=AF044311 /gj=3347841 /ug=Hs.63236 /en=708 | Cluster Ind L34155:Homo sapiens laminin- related protein (Lam.A3) mRNA, complete cds /cds=(0,5141) /gb=L34155 /gl=551596 /ug=Hs.83450 /Ren=5433 | Cluster Incl M69199:Human G0S2 protein gene, complete cds /cds=(160,471) /pb=M69199 /gi=609453 /ug=Hs.95910 /len=863 | Hs.174185 Cluster Incl L36594:Human autotaxin mRNA, complete cds (cds=(49,2798) (gb=13594 /gj=553905 /ug=Hs.174185 /en=3231 |
| Cluster# | Hs.347 | Hs.169470 | Hs.63236 | Hs.83450 | Hs.95910 | Hs.174185 |
| Gene Name | lactotransferrin | dystrophin (muscular dystrophy, Duchenne and Becker types), includes DXS142, DXS164, DXS206, DXS20, DXS239, DXS28, DXS288, DXS270, DXS272 | synuclein, gamma (breast cancer- specific protein 1) | laminin, alpha 3 (nicein (150kD), kalinin (165kD), BM600 (150kD), epilegrin) | putative lymphocyte G0/G1 switch gene | ectonucleotide pyrophosphatase/pho sphodiesterase 2 (autotaxin) |
| Genbank | U95626 | M18533 | AF044311 | L34155 | M69199 | L35594 |
| Affy | 37149_s_at | 40488_at | 36555_at | 37909_at | 38326_at | 41124_r_at |
| SeqID | 1168 | 1038 | 295 | 1023 | 1051 | 1024 |
| # | 136 | 137 | 138 | 139 | 140 | 141 |

| | 110 02/03/2 | -/1 | | 5 | 7 | | | 1,6502,021 | · |
|---------------------|---|--|--|--|---|---|--|---|--|
| p-values | 0.00041 | 1.65E-07 | 9.59E-07 | 6.44 E-0 7 | 0.00000343 | 0.0000486 | 0.000161 | 0.00000205 | 0.00000359 |
| Fold Change | 0.186645351 | 0.185170181 | 0.185137314 | 0.176679484 | 0.173326424 | 0.173040601 | 0.172026427 | 0.168054499 | 0.163077134 |
| Cluster Description | Cluster Incl. J02611:Human apolipoprotein D mRNA, complete cds /cds=(61,630) /gb=J02611 /gi=178840 /ug=Hs.75736 /fen=809 | Cluster Incl M91669:Hurnen Bullous pemphigoda autoartigen BP180 gene, 3 end Icds=(0,4598) /gb=M91669 /gi=179516 /lug=ls; 1/17938 /len=4669* | Cluster Incl X75958:H.sapiens trkB mRNA for protein-tyrosine kinase Iods=(97,1530) fg=-77958 (gi=-473007 /ug=+Rs.47860 fen=-2224 | Cluster Incl L35594:Human autotaxin mRNA, complete cds /cds=(49,2796) /gb=1,35594 /gi=537905 /ug=Hs.174185 /len=3231 | Hs.286116 Cluster Incl.L34041:Homo saplens L-glycerol-3 phosphate-NAD axidoreductase mRNA, compele cds Cds=(29,1078) (gb=L34041 /gl=508468 klig=Hs.28478 Ingn=1413 | L13740 /FEATURE= /DEFINITION=HUMTR3A Human TR3 orphan receptor mRNA, complete cds | Cluster Incl D45371:Human apM1 mRNA for CS3108 (novel adipose specific collagen-like factor), complete cds (cds=(26,780) (dp=D45371 /gi=87.1888 /ug=Hs.80485 Ilen=4517 | Cluster Incl AF-063002:Homo sapiens LIM protein SLIMMER MINAI, complete cds [cds=(84,1055) /gb=AF-063002 /gi=3859848 /ug=18-17629 /di=32042 | Cluster Incl X63741:H.sapiens pilot mRNA /cds=(353,1516) /gb=X63741 /gi=35472 /ug=Hs.74088 /len=4272 |
| Cluster# | Hs.75736 | Hs.117938 | Hs.47860 | Hs.174185 | Hs.286116 | Hs.1119 | Hs.80485 | Hs.239069 | Hs.74088 |
| Gene Name | apolipoprotein D | collagen, type XVII, alpha 1 | neurotrophic tyrosine kinase, receptor, type 2 | ectonucleotide pyrophosphatase/pho sphodiesterase 2 (autotaxin) | glycerol-3-phosphate dehydrogenase 1 (soluble) | nuclear receptor subfamily 4, group A, member 1 | adipose most abundant gene transcript 1 | four and a half LIM domains 1 | early growth response 3 |
| Genbank | J02611 | M91669 | X75958 | L35594 | L34041 | L13740 | D45371 | AF063002 | X63741 |
| Affy | 36681_at | 41618_at | 36042_at | 41123_s_at | 33902_at | 280 <u>g a</u> t | 40658_r_at | 32542_at | 40375_at |
| SedID | 1000 | 1056 | 1235 | 1024 | 1022 | 1016 | 996 | 301 | 1228 |
| #1 | 142 | 143 | 4 | 145 | 146 | 147 | 148 | 149 | 150 |

| | WO 02/0 | 59271 | | | 58 | | | PCT/U | S02/02176 |
|---------------------|--|--|--|---|-------------------|---|---|---|---|
| p-values | 0.00506 | 0.00282 | 0.0000162 | 5.79E-09 | 0.00000648 | 0.0000201 | 0.00000235 | 0.000768 | 0.0000385 |
| Fold Change | 0.161923599 | 0.161859881 | 0.155156674 | 0.145966282 | 0.143559713 | 0.138540058 | 0.137509192 | 0.137096706 | 0.136296847 |
| Cluster Description | Cluster Incl U15979:Human (dlk) mRNA, complete cds /cds=(173,1321) /gb=U15979 /gi=562105 /ug=Hs.169228 /len=1553 | K00650 /FEATURE=cds /DEFINITION=HUMFOS Human fos proto- oncogene (c-fos), complete cds | Cluster Ind X03350:Human mRNA for alcohol dehydrogensee beta-1-subunit (ADH1-2 allele) Icds=(72,1199) (gb=X03350 /gl=28415 AgH1-8,4 Men-2532 | Cluster Incl X64559:H.sapiens mRNA for tetranectin /cds=(93,701) /gb=X64559 /gj=37408 /ug=Hs.65424 /len=848 | | M57399 /FEATURE= /DEFINITION=HUMHBNF1 Human nerve growth factor (HBNF-1) mRNA, complete cds | Hs.274313 M62402 /FEATURE= /DEFINITION=HUMIGFBP4 Human insulin-like growth factor binding protein 6 (IGFBP8) mRNA, complete cds | Cluster Incl M12963;Human class i alcohol dehydrogenase (ADH1) alpha subunit mRNA, complete cds /cds=(72,1199) /gb=M12963 /gj=Hs,78089 /ug=Hs,78080 /ug=Hs,78048 Inn=1450 | i. Cluster Incl AB005293:Homo sapiens mRNA for perliptin, complete cus fcds=(124,1892) /gb=AB005293 /gi=3041770 /ug=Hs.103253 /ler=2904 |
| Cluster# | Hs.169228 | Hs.25647 | Hs.4 | Hs.65424 | Hs.74369 | Hs.44 | Hs.274313 | Hs.73843 | Hs.103253 |
| Gene Name | delta-like homolog (Drosophila) | v-fos FBJ murine osteosarcoma viral oncogene homolog | alcohol dehydrogenase 2 (class I), beta polypeptide | tetranectin (plasminogen-binding protein) | integrin, alpha 7 | pleiotrophin (heparin binding growth factor 8, neurite growth- promoting factor 1) | insulin-like growth factor binding protein 6 | alcohol dehydrogenase 1 (class I), alpha polypeptide | perilipin |
| Genbank | U15979 | K00650 | X03350 | X64559 | AF032108 | M57399 | M62402 | M12963 | AB005293 |
| Affy | 32648_at | 2094_s_at | 35730_at | 36569_at | 36892_at | 234_s_at | 1048 1736_at | 34637_f_at | 37122_at |
| SeqID | 1140 | 1009 | 1217 | 1229 | 293 | 1047 | 1048- | 1032 | 27.1 |
| ## | 151 | 152 | 153 | 154 | 155 | 156 | 157 | 158 | 92 |

| | , | WO 02/05 | 9271 | | | | PC | T/US02/02176 |
|---|---------------------|--|--|---|-------------------------------------|-------------------|---|---|
| | | | | | 5 | 9 | | |
| | sanja-d | 1.18E-0/ | 0.000082 | 0.0000134 | -9.26E-08 | 9.15E-08 | 0.000222 | 0.000439 |
| | Fold Change | 0.13521119 | 0.124249102 | 0.121953593 | 0.118530255 | 0.118243196 | 0.116969229 | 0.108941232 |
| | Cluster Description | Cluster Incl AJ001183:Homo septens mRNA for Sox10 protein (cds=(120.1520) fp=AJ001183 (gi=2909359 /ug=Hs.44317 | Inen-2x9. Classer Incl L49169:Human G0S3 mRNA, complete cds /cds=(593,1609) /gb=L49169 /gi=1082037 /ug=Hs.75678 /len=5775 | A Human CIT987SK-A- | | | 50 M21829 FEATURE=mRNA DEFINITION=HUMRERZA Human keratin type II (58 kD) mRNA, complete cds | Cluster Incl AA128249;zi29d09.r1 Homo sapiens cDNA, 5 end /clone=IMAGE-603345 /clone_end=5" (gp=AA128249 /gi=1688343 /ug=Hs.83213 /len=648" |
| ì | Cluster# | Hs.44317 | Hs.75678 | Hs.78344 | Hs.155597 | Hs.283750 | Hs.195850 | Hs.83213 |
| | Gene Name | SRY (sex determining region Y)-box 10 | FBJ murine osteosarcoma viral oncogene homolog B | myosin, heavy polypeptide 11, smooth muscle | D component of complement (adipsin) | tenascin XA | keratin 5 (epidermolysis bullosa simplex, Dowling- Mearaf/Kobner/Weber- | cockayne types) fatty acid binding protein 4, adipocyte |
| | Genbank | AJ001183 | L49169 | AF001548 | M84526 | U89337 | M21389 | AA128249 |
| | Affy | 36018_at | 36669_at | 767_at | 40282_s_at | 38508_s_at U89337 | 613_at | 38430_at |
| | Glbes | 849 | 1029 | 283 | 1055 | 1166 | 1039 | 45 |

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| | | | | | ••• | | | |
|---------------------|---|--|--|---|--|--|---|--|
| p-values | 9.35E-07 | 5.83E-07 | 0.000249 | 0.00000084 | 0.0000236 | 2.66E-08 | 3.51E-07 | 0.00000596 |
| Fold Change | 0.108244989 | 0.10121597 | 0.097312672 | 0.096634153 | 0.092826583 | 0.091111614 | 0.090848213 | 0.070781449 |
| Cluster Description | Cluster Incl U95367:Human GABA-A receptor pi subunit mRNA, complete cds cds=(156,1478), {gb=U95367 /gi=2197000 /lig=F,70725 /len=3284 | Cluster Incl M57399:Human nerve growth factor (HBNF-1) mRNA, complete cds cds=295,901) /gb=M57399 /gi=292072 /ig=Hs.44 /len=1029 | Cluster Incl X07696:Human mRNA for cytokeratin 15 /cds=(61,1431) /gb=X07696 /gj=34070 /ug=Hs.80342 /len=1709 | Cluster Incl M69225:Human bullous pemphigoid antigen (BPAG1) mRNA, complete ods /ods=UNKNOWN /gb=M69225 /gi=179522 /ug=15.620 /len=9830 | Cluster Incl AF013570:Homo sapiens smooth muscle myosin heavy chain SM2 mRNA, alternatively spliced, partial cds /cds=(0,1767) /gb=AF013570 /gj=2352944 /ug=Hs.78344 /hen=2880 | Cluster Incl D17408:Homo sapiens mRNA for calponin, complete ods kots=(92,985) (gb=17408 (gi=1783204 /ug=Hs.21223 /en=1517 | D00654 /FEATURE=cds //DEFINITION=HUMACTSG7 Homo sapiens gene for enteric smooth muscle gamma-actin, exons, complete cds | Cluster Incl. J00124:Homo sepiens 50 kDa type I epidemmal keratin gene, complete cds /cds=(61,1479) (gp=J00124 /gi=186704 /ug=Hs.117729 /len=1634 |
| Cluster # | Hs.70725 | Hs.44 | Hs.80342 | Hs.620 | Hs.78344 | Hs.21223 | Hs.77443 | Hs.117729 |
| Gene Name | gamma-aminobutyric acid (GABA) A receptor, pi | plelotrophin (heparin binding growth factor 8, neurite growth- promoting factor 1) | Keratin 15 | bullous pemphigoid antigen 1 (230/240kD) | myosin, heavy polypeptide 11, smooth muscle | calponin 1, basic, smooth muscle | actin, gamma 2, smooth muscle, enteric | keratin 14 (epidermolysis bullosa simplex, Dowling-Meara, Koebner) |
| Genbank | U95367 | M57399 | X07696 | M69225 | AF013570 | D17408 | D00654 | J00124 |
| Affy | 40339_at | 34820_at | 37582_at | 40304_at | 37407_s_at | 34203_at | 1197_at | 39052_at |
| SedID | 1167 | 1047 | 1220 | 1052 | 288 | 964 | 696 | 866 |
| # | 167 | 168 | 169 | 170 | 171 | 172 | 173 | 174 |
| | | | | | | | | |

| | | | 61 | | | |
|---|---|--|---|--|---|--|
| p-values 5.77E-09 | 0.00055334 | 0.00136502 | 4.5016E-06 | 0.00129495 | 0.00039175 | 2.7406E-06 |
| Fold Change 0.0564074 | 7.73694268 | 4.18 | 0.282343722 | 3.871753195 | 0.308362018 | 0.227899817 |
| Cluster Ind AF056087 Homos aspiens secreted firzated related protein mRNA, compilete dis lods=(302,1243) /gb=AF056087 /gj=3033550 /ug=Hs.7306 /her=4488 | Cluster Ind. AW007442:wt55g06.x1 Homo sapiens cDNA, 3 end /clone=IMAGE-2511418 /clone_end=3 /gb=AW007442 /gj=5856220 /ug=18_235651 /len=424 | | Cluster Incl. AA614135:no8209.s1 Homo sapiens cDNA, 3 end /donne-IIMAGE-1113353 /clone_end=3'/gb-A614135 /gl=2466269 /tuc+16,49765 /fne=664' | | Hs.188961 Cluster Ind. AI283888:qk51f12.x1 Homo sapiens cDNA, 3 end /done=INIA/GE-1872527 /done_end=3/gb=AI283888 /gj=3922.121 /done=is-153544 /len=420 | Cluster Incl. AI599804;tm7/e11.x1 Homo sapiens cDNA, 3 end /done=IMAGE-2163884 /clone_end=3' /gb=AI598904 /gi=4598852 /ug=Hs.12828 /len=508' |
| Cluster# | Hs.821 | Hs.179718 | Hs.49765 | Hs.151678 | Hs.186961 | Hs.12828 |
| Gene Name secreted frizzled- related protein 1 | zinc finger protein homologous to Zfp92 in mouse | v-myb avian myeloblastosis viral oncogene homolog- like 2 | VLCS-H1 protein | UDP-N-acetyl-alpha-D-galactosamine:polype ptide N-acetylgalactosaminyltr acetylgalactosaminyltr ansferase 6 (Gall No-T6) | ubiquitin specific protease 25 | tweety (Drosophila) homolog 1 |
| Genbank AF056087 | AW007442 | A1990026 | AA614135 | AL118633 | Al283888 | Al589804 |
| Affy 32521_at | 74989_at | 80675_at | 64695_at | 59253_at | 91419_at | 48513_at |
| SeqID 299 | 923 | 843 | 183 | 901 | 411 | 525 |
| 175 | 176 | 177 | 178 | 179 | 180 | 18 |

-

| | DCT/HS02/02176 | |
|---|--|--|
| 0.00022486 | 6.75E-04 | |
| 0.149624624 0.00022486 | 4.122293677 | |
| /ug=rs.14319 /len=52/ Cluster Incl. Al480357:tm51d08.x1 Homo sapiens cDNA. 3 end /clone=IMAGE-2161647 | /done_and=3' igb=A4480357 /gi=4373525 /ug=1% s177 /aim=330 /2536 Cluster Incl A4704137/ag47g01 s1 Homo saptiens cDNA, 3 and /done=IMAGE-1119894 /clone_and=3" (gb=A4704137 /gi=2714055 /ug=1% 125359 fan=923" | |
| 1877 | 2535 | |

| | WO 02/059 | 271 | | 6 | 2 | | PC | T/US02/02176 |
|---------------------|---|--------------------|---|----------------------|---|---|--|-------------------------------|
| p-values | 0.00017707 | 1.0421E-05 | 3.9817E-06 | 1,5585E-05 | 7.4333E-10 | 0.00109754 | 0.00022486 | 6.75E-04 |
| Fold Change | 6.818460543 | 4.363660627 | 4.2443289 | 4.256819836 | 9.431588747 | 5.50 | 0.149624624 | 4.122293677 |
| Cluster Description | Hs.159154 Cluster Incl. AW003626:wx34e02.x1 Homo sapiens cJNA, 3 end (done=IMAGE-2545652 /done_end=3' (gb=AW003628 /gl=5850542 /ug=ls.224018 /len=707 | | Hs.104019 Ctuster Incl. A1990842.ws22g06.x1 Homo saplens cDNA, 3 end /dome=1/AAGE-2487978 /done_end=3 /gb=A1990842 /gi=5837523 /uc=Hs.233896 /fen=591 | | Cluster Incl. AW003286:wq65a02.x1 Homo sapiens cDN4, 3 end (clone=IMAGE-2476106 fclone_end=3' (gb=AW003286 /gj=5850202 fuce+1s_232126 fnen=773' | Cluster Incl. AA707213:zj32h06.s1 Homo sapiens cDNA, 3 end (clone=452027 lclone_end=3' (gb=AA707213 /gj=2717131 /luc=16: 14319 /len=527 | Cluster Incl. Al480357:tm51d08.x1 Homo sapiens cDN4, 3 end /done=IMAGE-21d1647 /done_end=3 /gb-4480357 /gi=4373525 /uc=16.5 g1877 /len=390 | |
| Cluster# | Hs.159154 | Hs.169840 | Hs.104019 | Hs.108106 | Hs.270810 | Hs.317 | Hs.91877 | Hs.125359 |
| Gene Name | tubulin, beta, 4 | TTK protein kinase | transforming, acidic coiled-coil containing protein 3 | transcription factor | topolsomerase (DNA) Hs.270810 II alpha (170kD) | topoisomerase (DNA) I | thyroid hormone responsive SPOT14 (rat) homolog | Thy-1 cell surface antigen |
| Genbank | AW003626 | AI973225 | A1990642 | AA026429 | AW003286 | AA707213 | AI480357 | AA704137 |
| Affv | 9200 | 74593_at | 52238_s_at Al990642 | 63346_at | 74096_at | 90442_at | 57778_at | 39395_at |
| SealD | 913 | 833 | 847 | 9 | 911 | 208 | 488 | 205 |
| # | 182 | 83 | 184 | 185 | 186 | 187 | 188 | 189 |

| | PCT | /US0: | 2/0217 | 6 |
|---|--|---|--|------------------------|
| 0.00133643 | | 0.0000138 | | |
| 0.325281828 0.00133643 | | 0.277326129 0.0000138 | | |
| /ug=rrs.zzuos /ret i=oou Hs.24395 Cluster Incl. N45415;yw97h06.r1 Homo | sapiens cDNA, 5 end /done=IMAGE-260219 /done_end=5'/gb=N45415 /gi=1186581 | /ug=Hs.24395 /len=64 / Hs.20144 Cluster Ind Al720438:as81g04.x1 Homo | sapiens cDNA, 3 end /clone=IMAGE-2335158 /clone_end=3" /gb=AI720438 /gi=5037694 | /ug=Hs.20144 /len=550" |
| Hs.24395 | | Hs.20144 | | |
| small inducible | cytokine subfamily B (Cys-X-Cys), member | 14 (BRAK) small inducible | cytokine subfamily A (Cys-Cys), member | 14 |

| | WO 02/059 | 271 | | 63 | | | PCT/U | IS02/02176 |
|---------------------|--|--|--|---|---|---|---|--|
| p-values | 0.00065669 | 3.1699E-09 | 0.00058351 | 3.4432E-06 | 2,6069E-05 | 0.00051209 | 0.00133643 | 0.0000138 |
| Fold Change | 3.418467862 | 0.284181885 | 99.9 | 0.31 | 4.206790316 | 3.273076721 | 0.325281828 | 0.277326129 |
| Cluster Description | Cluster Incl. A4527151:ni07b08.s1 Homo sepiens cDN4, 3 end (clone=IMA/GE-967287 dlone_end=3'/gb=A4527151 /gi=2269220 /lig=416.108977 Men=559 | | Cluster Incl. AL036753:DKFZp564l0663_r1 Homo sapleins cDN4, 5 end Hone=DKFZp564l0663 (Abne_end=5' fgb=AL036753 /gl=5927893 /ug=Hs.133482 Aen=617 | sparm surface protein Hs.129872 Cluster Incl. Al949433:wq11e08.x1 Homo saplens 6D/NA, 3 end vlaone-liNAGE-2470982 clane_and=2470982 clane_and=37 (pj=Al940433) gj=5741831 / lug+Hs.13962 len=661* | Cluster Ind. AI285531:qu49b04.x1 Homo sapiens cDN4, 3 end /dolone=IMAGE-1988079 /done_end=3' /gb=AI285531 /gi=3923764 /uq=Hs. 106260 /len=598 | Cluster Incl. H10816;ym04e12.r1 Homo saptens cDNA, 5 end /clone=INAGE-46684 iclone_end=5' /gb=H10816 /gi=875636 /ug=Hs.22065 /fen=660' | Cluster Incl. N45415;yw97h06.r1 Homo sapiens cDNA, 5 end /done=INAGE:260219 /done_end=5 /gb=N45415 /gl=1186581 /ug=18,24395 /len=647 | Cluster Ind Al720438:as81g04.x1 Homo sapiens cDNA, 3 end /donn=IMAGE-2355158 /folone_and-3" (pa 4720438 /gi=5037694 /incl+6 70144 liben=FR)" |
| Cluster # | Hs.184242 | Hs.243678 | Hs.32964 | Hs.129872 | Hs.106260 | Hs.22065 | Hs.24395 | Hs.20144 |
| Gene Name | sterol-C5-desaturase (fungal ERG3, delta-5 desaturase)-like | SRY (sex determining Hs.243678 region Y)-box 8 | SRY (sex determining region Y)-box 11 | sperm surface protein | sorting nexin 10 | small protein effector 1 of Cdc42 | small inducible cytokine subfamily B (Cys-X-Cys), member 14 (BRAK) | small inducible cytokine subfamily A (Cys-Cys), member |
| Genbank | AA527151 | AI675177 | AL036753 | Al949433 | Al285531 | H10816 | N45415 | AI720438 |
| Affy | 48101_at | 43039_at | 91880_at | 78444_at | 63335_at | 49498_at · H10816 | 65823_at | 33790_at |
| SealD | 152 | 285 | 854 | 800 | 412 | 980 | 1077 | 616 |
| ## | 190 | 191 | 192 | 193 | 194 | 195 | 196 | 197 |

| WO 0 | 2/059271 | | | 64 | | PC1/USC | 12/02176 |
|--|--|---|---|-----------------|--|--|----------------------------|
| p-values 1.7128E-08 | 0.000166 | 5.5177E-07 | 2.1786E-06 | 0.02422274 | 0.00029667 | 3.31E-03 | 8.0344E-05 |
| Fold Change 0.13254339 | 0.147972137 | 0.279118305 | 0.113932898 | 3.52 | 5.371729632 | 3,31996447 | 5.815956961 |
| Cluster Description Cluster Incl. AA102575-zn42c03.s1 Homo sapiens cDNA, 3 end /clone=INAGE-550084 | Iclone_end=5' (pl=A41025/5 /gl=1841/6/ I/ug=Hs.26530 (len=689' Cluster Incl A4829286:cr08801.s1 Homo sapiens cDNA, 3 end (done=IMAGE-1420488 -in-lone and=3'' (h=A8P09786 /n=207385) | | MgH8.2380 Inflants All QFH8.2380 Inflants Cluster Inc. Al829365:wk64e08.x1 Homo sapiens cDNA, 3 end /done=IMAGE-2420198 All Clone, end-3' (pla-pl829385 /gl=5450056 | | Cluster incl. AIR6401 6xwj53hrt0.xr Homo saplens Collay, 3 and Alchore=IIANGE-2406895 Adone and=3' tgb=AIR64016 (gj=5528123 hug=Hs.234375 /len=5111 | Cluster Incl AA131149:zo16d05.r1 Homo sapiens cDNA, 5 end /done=INAGE-587049 /done_end=5" /gb=4A131149 /gi=1692640 /uc=Hs.2962 len=464" | |
| Cluster# Hs.26530 | Hs.181062 | Hs.1327.92 | Hs.59729 | Hs.279623 | Hs.313 | Hs.2962 | Hs.194691 |
| Gene Name serum deprivation response | (phosphatidylserine- binding protein) serum arryloid A1 | serologically defined colon cancer antigen 43 | semaphorin sem2 | selenoprotein X | secreted phosphoprotein 1 (osteopontin, bone staloprotein I, early T- lymphocyte activation | S100 calcium-binding protein P | retinoic acid induced 3 |
| Genbank AA102575 | AA829286 | AI832477 | AI829385 | W92110 | Al864016 | AA131149 | A1990405 |
| Affy 50094_at | 33272_at | 63580_at | 62486_at | 78757_at | 74815_at | 34319_at | 57027_at |
| SeqID 39 | 233 | 732 | 729 | 1213 | 746 | 46 | 844 |
| 198 | 199 | 200 | 201 | 202 | 203 | 204 | 202 |
| | | | | | | | |

| WO 02/059271 | | | 65 | | | PCT/US0 | 2/02176 |
|--|--|--|--|---|--|--|--|
| p-values 4,5657E-06 | 0.02812425 | 0.00031747 | 4.4105E-06 | 2.9782E-08 | 1.3574E-05 | 2.9836E-06 | 9.3518E-10 |
| Fold Change 0.18 | 0:30 | 3.28 | 4.031107597 | 6.276196882 | 0.26701338 | 0.183592799 | 5.975488989 |
| Cluster Description Cluster Incl. At 110389.DNFZp564P1078_s1 Homo saplens GNM, 3 end Idono=DNFZp564P1078 Iclone_end=3** Igo=Att.10388 Igi=5868976 Iugi=18:188383 Illon=Tio | Cluster Incl. AA147088:zo32c07.s1 Homo sapiens cDNA, 3 end /clone=IMAGE-588588 clone_end=3'/gp=AA147088 /gl=1716461 /lig=ls,106185 /len=688 | Cluster Ind. Al799107:we98h02.x/1 Homo sepiens cDNA, 3 end /clone=IMAGE-2349171 /clone_end=3'/gb=Al799107 /gi=5364579 /tg=518501 /len=713' | Cluster Inci. AL079372:DKFZp564H1178_s1 Homo sapleirs cDN4, 3 end Idone=DKFZp684H1178 Idone_end=3' Igb=AL078372 (gi=5423266 /ug=Hs_23044 Iden=555' | Cluster Incl. AI739117:wi18c04.x1 Homo sapiens cDN4, 3 end /clone=INAGE-2300598 /clone_end=3'/gb-A739117/gl=5101098 /tuc=16,73525 /fen=762' | Cluster Incl. Al858625:w440g05.x1 Homo sapiens cDNA, 3 end /done=INAGE-2427416 /done_end=3' /gb-B58626 /gi=5512242 /ug=Hs.30807 /len=617 | Cluster Incl. Al304339:qo58g10.x1 Homo saplens cDNA, 3 end /done=INA/GE-1912770 /done_end=3 /gb-Al304339 /gi=3988028 /ug=Hs.8364 /len=514* | Cluster Incl. AA195614:zr37b09.r1 Homo sapiens cDNA, 5 end /done=IMA/GE-685561 /done_end=5 /gb=AA195614 /gi=1783791 /uc=Hs.5/g1 /len=535 |
| Cluster# Hs.12246 | Hs.106185 | Hs.128501 | Hs.23044 | Hs.73625 | Hs.198281 | Hs.8364 | Hs.5101 |
| Gene Name reelin | ral guanine nucleotide Hs.106185 dissociation stimulator | RAD54, S. cerevisiae, Hs.128501 homolog of, B | RAD51 (S. cerevisiae) homolog (E coli RecA homolog) | RAB6 interacting, kinesin-like (rabkinesin6) | pyruvate kinase, muscle | pyruvate dehydrogenase kinase, isoenzyme 4 | protein regulator of cytokinesis 1 |
| Genbank AL110368 | AA147088 | AI799107 | AL079372 | AI739117 | AI858626 | Al304339 | AA195614 |
| Affy 69473_r_at | 91384_at | 74300_at | 44037_at | 46683_at | 57173_at | 48647_at | 45799_at |
| SeqID 900 | 2 | 22.0 | 895 | 620 | 737 | 420 | 20 |
| 50e | 207 | 208 | 506 | 210 | 211 | 212 | 213 |
| | | | | | | | |

| | | | | | 56 | | | |
|---------------------|---|---|---------------------|--|---|---|--|---|
| p-values | 1.0797E-05 | 5.5208E-05 | | 0.00261362 | 7.8961E-05 | 0.00044405 | 9.7327E-07 | 6.0939E-06 |
| Fold Change | 0.31 | 0.181794989 | 3.638706185 | 0.158415546 | 0.062026919 | 0.19 | 0.14 | 5.25 |
| Cluster Description | Cluster Incl. AI828396:wk34e07.x1 Homo septiens coltnk, 3 end (done=IMAGE-2422.116 Idone_end=3' (gb-AI82396 (gj=5449067 hg=1s.201574 Hen-466 | Cluster Incl. AI741776:wg22g10.x1 Homo sapiens CDNA, 3 end /done=IMAGE-2365890 /done_end=3' /gb=AI741776 /gj=5110064 /ug=les.8944 /len=641 | | Cluster Incl. Al863965:wj54508.x7 Homo saplens CDNA, 3 end /done=IMAGE-2406615 /done_end=3' /gb=Al863965 /gi=5527996 /lig=16, 15295 /len=611' | | Cluster Incl. AL044906:DKFZp434K183_r1 Homo saplens cDN4, 5 end Iclone=DKFZp434K183 (done_end=5' fgb=AL044906 (gi=5433103 /ug=Hs.211647 Ilen=485' | Cluster Incl. Al492388:tt27d10.x1 Homo sapiens cDNk, 3 end /done=INAGE:-2131699 /clone_end=3 /gb=Al492388 /gi=4393391 /lug=h.145011 /len=477 | Cluster Incl. AA536819:nj/79e01.sr Homo sapiens cDNA klone=IMAGE-998712 /gb=A4536819 /gj=2280072 lug=Hs.238355 /en=522 |
| Cluster # | Hs.75323 | Hs.8944 | Hs.163833 | Hs.279898 | Hs.205126 | Hs.75813 | Hs.117176 | Hs.83883 |
| Gene Name | prohibitin | procollagen C- endopeptidase enhancer 2 | PRO0611 protein | PRO0529 protein | polymeric immunoglobulin receptor | polycystic kidney disease 1 (autosomal dominant) | poly(A)-binding protein, nuclear 1 | PMEPA1 protein |
| Genbank | AI828396 | AI741776 | AA593830 | AI863965 | Al697470 | AL044906 | Al492388 | AA535819 |
| Affy | 86573_at | 55569_at | 84574_r_at AA593830 | 65700_at | 45294_at | 71106 <u> </u> at | 90494_at | 91095_s_at |
| SeqID | 726 | 625 | 173 | 745 | 607 | 874 | 492 | 161 |
| #1 | 214 | 215 | 216 | 217 | 218 | 219 | 220 | 221 |
| | | | | | | | | |

| WO 02/033 | .2/1 | | • | 7 | | 10 | 1/0502/021/0 |
|--|---|------------------------------|--|--|--|--|--|
| p-values 5.9471E-05 | 0.00024055 | 5.6196E-05 | 0.00023065 | 0.00627388 | 1,9181E-06 | 2.4915E-05 | 2.4435E-05 |
| Fold Change 0.325587729 | 0.224464715 | 0,285584964 | 0.257023901 | 0.28 | 0.133294383 | 0.25 | 0.29 |
| Cluster Incl. ABY1679 wq88b01.x1 Homo Cluster Incl. ABY1679 wq88b01.x1 Homo Belsens CDMA, 3 and Johne=IMAGE-2479081 Iclone_end=3' (gp=AI971679 (gl=5768505 Iug=Hs.239374 Jen=573 | Cluster Incl. AI815028:wk70b07.x1 Homo sapiens cDNA, 3 end /clone=IMAGE-2420725 /clone_end=3 /gb-AI815028 /gj=5426243 /ug=Hs.5228 /lan=492* | | Cluster Incl. AA521373:aa77g02.s1 Homo sapiens cDNA, 3 end /clone=IMAGE-828994 /clone_end=3'/gb=AA521373 /gi=2261916 /ug=Hs,9469 /Ren=526 | Cluster Incl. AW/007686:wt02e03.x1 Homo saplens clothk, 3 end /clone=IMA/GE-2506300 /clone_end=3'/gb=AW/007566 /gi=5856429 /ug=Hs.229193 (Ren=52.4 | Cluster Incl. T66157.yc77d05.s1 Homo sapiens 0.133294383 cDNA, 3 end /donne=IIMAGE-22165 /done=IIMAGE-22165 /done=IIMAGE-22165 /done=IIMAGE-22165 /done=IIMAGE-23001 /done-416 | | Oluster Incl. AI851185;vx64e12;x1 Homo sapiens cDNA, 3 end /clone=IMAGE-2548462 /clone_end=3'/gb=AI951185 /gl=5743495 /ug=Hs:144630 /len=787 |
| Cluster# Hs.75825 | Hs.82101 | Hs.182538 | Hs.9469 | Hs.1872 | Hs.154437 | Hs.173560 | Hs.144630 |
| Gene Name pleiomorphic adenoma gene-like 1 | pleckstrin homology- like domain, family A, member 1 | phospholipid scramblase 4 | phospholnositol 3- phosphate binding protein-1 | phosphoenolpyruvate carboxykinase 1 (soluble) | phosphodiesterase 2A, cGMP-stimulated | odd Oz/fen-m homolog 2 (Drosophila, mouse) | nuclear receptor subfamily 2, group F, member 1 |
| Genbank Al971679 | AI815028 | AI818248 | AA521373 | AW007566 | T66157 | AI859144 | Al951185 |
| Affy 57266_r_at | 62196_at | 59010_at | 63017_at | 80604_at | 90033_at | 77546_at | 78518_at |
| 40 | w | | | | | | |
| SeqID 824 5 | 869 | 705 | 143 | 924 | 1128 | 738 | 804 |
| | | 224 705 | 225 143 | 226 924 | 227 1128 | 228 738 | 229 804 |

| WO 02/059271 | | | 68 | | | PC1/US02/021/6 | | |
|--------------|--|--|--------------------------------|--|--------------------------------------|---|--|--|
| | p-values 0.00020188 | 0.00307749 | 6.6167E-05 | 0.000479 | 1.4629E-08 | 6.7348E-06 | 4.93E-06 | 5.3066E-07 |
| | Fold Change 0.199980315 | 3.316200333 | 4.64 | 0.117240353 | 0.17504844 | 0.262530591 | 4.698983711 | 3.809930405 |
| | Cluster Description Cluster Ind. A136848;qu020403.r1 Homo saplens cDNA, 3 end (Abone=IMAGE-1963601 (Abone_end=3' (gp=A1356848 /gj=4096001 //ug=Hs.239464 /en=717 | Cluster Incl. Al935353.wo82d12.x1 Homo sapiens cDN4, 3 end /clone=IMAGE-2461847 /clone_end=3' /gb=Al935353 /gi=5674223 /uc=ls.234669 /len=532" | | Cluster Ind. W48800.zc44f03-r1 Homo sapiens 0.117240353 cluxe, ind. W48800.zc44f03-r1 Homo sapiens 0.117240353 clow, 5 end clone=liMAGE-325f81 forme_and=3 /gb=W48800 /gi=1336949 /ineHs. if7297 /len45f97 | | Cluster Incl. A1052524:oz27f07.x1 Homo saplens cDN4, 3 end /done=IMAGE-1676581 /done_end=3' /gb=A1052524 /gi=3308515 /incl+6 4700 /lone_537 | Cluster Incl AA4/18080zx97h07.s1 Homo stephers DA7A, 3 and Clong=III/AA;E;-767773 Clone, end-s²' týp=AA4/18080 (gj=2079881 /tg=Hs.3972 /len=54.3" | Cluster Incl. AI742260:wg39g10.x1 Homo sapiens cDNA, 3 end /clone=IMAGE-2367522 /clone_end=3' /gb=AI742260 /gl=5110548 /ug=Hs.93847 /len=789 |
| | Cluster# | Hs.243886 | Hs.159623 | Hs.44054 | Hs.266902 | Hs.3821 | Hs.3972 | Hs.93847 |
| | Gene Name nuclear factor I/X (CCAAT-binding transcription factor) | nuclear autoantigenic Hs.243886 sperm protein (histone-binding) | NK-2 (Drosophila) homolog B | ninein | neurotrophin 5 (neurotrophin 4/5) | neurobeachin | NeuAc-alpha-2,3-Gal- beta-1,3-GalNAc- alpha-2, 6- sialyltransferase alpha2,6- sialytransferase | NADPH oxidase 4 |
| | Genbank Al355848 | Al935353 | AI479933 | W48800 | AA427578 | AI052524 | AA418080 | Al742260 |
| | <u>Affy</u> 63877at | 49666_s_at | 72236_at | 82657_f_at | 72026_g_at AA427578 | 56809_at | 34778_at | 55457_at |
| | SeqID 435 | 790 | 486 | 1188 | 1 4 | 328 | 105 | 630 |
| | 230 # | 231 | 232 | 233 | 234 | 235 | 236 | 237 |

| | | | | 69 | | | | |
|---------------------|-------------------------------------|--|--|--|---|---|--|--|
| p-values | 4.4362E-07 | 0.0000139 | 0.00225117 | 0.00151003 | 1.1301E-05 | 0.0000045 | 0.00267462 | 0.0000502 |
| Fold Change | 0.32670528 | 0.168769332 | 0.288946314 | 4.848524211 | 3.068295724 | 0.23811529 | 0.23 | 0.246437105 |
| Cluster Description | | Cluster Incl AF001548:Human Chromosome If BAC done CIT9875K-A-815A9 (cds=(0,5573) (gb=AF001548 /gi=2104552 h(g=ls,78344 /len=6428 | Cluster Inc., Al814178xw62s10.x1. Homo suphers Colva, 8 and Idone=IMAGE-2420106 Ichone, and-8' (gp-AR1718 (gp-E425393 Iug=Hs.234582 /hen=408' | Cluster Incl. N78139;yy78d07.r1 Homo sapiens 4,848524211 cDNA_5 ein clothor=INAGE-248865 clothe_end=5/gb=N78139 (gi=1240840 lug=Hs.239825 flen=656 | Cluster Incl. N27428;yx81th09.s1 Homo sapiens CDNA, 3 end (donne=IMAGE-288193 fdone_end=3'/gb=N27428 (gi=1141909 fug=18.221178 Hen=407 | Clusier Incl AA420624nc61c12.r1 Homo sapiens cDN4 /clone=INAGE-745750 /dp=AA426624 /gi=2094502 /ug=Hs.183109 /fen=533 | Cluster Ind. AA776393:ah16g09.srl Homo sapiens ODNA, 3 and /dolne=1158884 //done_end=3'/gb=AA776393 /gj=2835727 //g=18.177536 /len=602' | Cluster Incl AA461385.zx70e07.r1 Homo sapiens cDNA, 5 end /clone=IMAGE-796836 /clone_end=5" /gb=AA461365 /gl=2186485 /ug=Hs.237742 /len=591" |
| Cluster # | Hs.211582 | Hs.78344 | Hs.239663 | Hs.42346 | Hs.240 | Hs.183109 | Hs.177536 | Hs.279771 |
| Gene Name | myosin, light polypeptide Kinase | myosin, heavy polypeptide 11, smooth muscle | myeloid/lymphoid or mixed-lineage leukemia (trithorax (Drosophila) homolog); translocated to, 7 | muscle-specific protein | M-phase phosphoprotein 1 | monoamine oxidase A | metallocarboxypeptid ase CPX-1 | melanoma inhibitory activity,ras-related GTP-binding protein 4b |
| Genbank | AA526844 | AF001548 | Al814178 | N78139 | N27428 | AA420624 | AA776393 | AA461365 |
| Affy | 46276_at | 32582_at | 46743_s_at | 58494 <u>r</u> at | 59624_g_at N27428 | 41771_g_at AA420624 | 84934_at | 39271_at |
| SeqD | 150 | 283 | 269 | 1093 | 1069 | 111 | 222 | 127 |
| # | 238 | 239 | 240 | 241 | 242 | 243 | 244 | 245 |

| 70 | | | | | | | | |
|---------------------|--|--|--|---|--------------------------------------|--|--------------------------------|-----------------------------|
| p-values | 6.1415E-06 | 1.9615E-06 | 0.00253759 | 0.00110724 | 0.0005914 | 0.00014885 | 8.7304E-05 | 6.1228E-07 |
| Fold Change | 0.196704294 | 0.12 | 0.307354772 | 4.366999247 | 0.247555921 | 0.214230439 | 0.322163154 | 3.542386026 |
| Cluster Description | Cluster Incl. AB72357:wr37c09.x1 Homo sapiens GolNA, 3 end (Jone=IMAGE-2489872 I/done_end=3' (gb=A1972357 /gi=5769183 /ug=H5.257874 Ien=524' | Cluster Incl. Al459139:ij65e05.x1 Homo sapiens cDNA, 3 end clone=INA/GE-2146400 fidone_end=3' (gb=Al459139 /gi=4311718 /ug=Hs.235500 /len=519' | Cluster Incl. Al969879:wq76f05.x1 Homo sapiens cDNA, 3 end (done=IMAGE-2477217 Idone_end=3' (gb=Al969879 /gj=5786897 /ug=Hs.233610 /len=329' | Cluster Incl. W22091:61F10 Homo sapiens cDNA /dione=(not-directional) /gb=W22091 /gi=1298924 /ug=Hs.234460 /len=637 | | Cluster Ind. Al815057:wk70e06.x1 Homo saplens cDNA, 3 end /clone=IMAGE-2420770 /done_end=3' /gb=Al815057 /gj=5426272 /ug=Hs.88559 /fen=484' | | |
| Cluster # | Hs.19368 | Hs.56729 | Hs.89137 | Hs.1770 | Hs.97044 | Hs.84728 | Hs.182965 | Hs.279766 |
| Gene Name | matrilin 2 | lymphocyte-specific protein 1 | low density lipoprotein Hs.89137 related protein 1 (alpha-2- macroglobulin receptor) | ligase I, DNA, ATP- dependent | Kv channel- interacting protein 2 | Kruppel-like factor 5 (intestinal) | Kruppel-like factor 4 (gut) | kinesin family member 4A |
| Genbank | Al972357 | Al459139 | Al969879 | W22091 | F36908 | AI815057 | AI290876 | AA004208 |
| Affy | 75254_at | 78644_at | 74641_at | 74835_r_at | 48950_at | 47113_at | 48587_at | 51160_at |
| SedD | 829 | 480 | 819 | 1176 | 976 | 669 | 416 | 2 |
| # | 246 | 247 | 248 | 249 | 250 | 251 | 252 | 253 |

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|---|---------------------|---|------------------|------------------|---|--|---|---|---|
| | sanjev-d | 3.2252E-06 | 0.00090706 | 0.00048007 | 0.00894957 | 9.936E-05 | 4.1926E-05 | 1.24E-12 | 1.69E-06 |
| | Fold Change | 0.189411148 | 0.317853152 | 0.28929478 | 3.778012983 | 0.231260722 | 0.20 | 10.18941323 | 3.635450964 |
| | Cluster Description | Cluster Incl. W68504.zd36b01.r1 Homo sapiens cDNA, 5 end /clone=IMAGE-342697 clone_end=5′ (gb=W68504 /gj=1377374 /tig=1s_27298 /fen=613° | | | Cluster Incl. AI073544:0v45e02.x1 Homo sapiens cDNA, 3 end /clone=IMAGE-1640282 /clone_end=2' /gb=AI073544 /gi=3400188 /ug=Hs.120703 /len=481* | Cluster Ind. Af097463:qb90e04.x1 Homo seplens cDN4, 3 end /done=IMAGE-1707342 /done_end=3'/gb=Al097463/gi=3447045 /ug=18.2/1035/fen=464' | Cluster Incl. H19400;ym48g07.r/ Homo sepiens cDNA, 5 end /clone=IMAGE-51511 /clone_end=5/ gb=H19400 (gl=888095 /tg=Hs,2r1035 /fen=488' | Cluster Incl AB029000:Homo sepiens mRNA for KIAA1077 protein, partial cds /cds=(0,2456) /gb-AB029000 /gi=5689490 /ug=Hs,70823 /len=4834 | Cluster Incl AB020713:Homo sapiens mRNA for KIAA0906 protein, partial cds /cds=(0,2772) /gb=AB020713 /gi=4240300 /ug=Hs.56966 |
| | Cluster # | Hs.191098 | Hs.230188 | Hs.263395 | Hs.62576 | Hs.21035 | Hs.21035 | Hs.70823 | Hs.56966 |
| : | Gene Name | KIAA1479 protein | KIAA1396 protein | KIAA1368 protein | KIAA1240 protein | KlAA1130 protein | KIAA1130 protein | KIAA1077 protein | KIAA0906 protein |
| | Genbank | W68504 | AA 912409 | AF063500 | A1073544 | Al097463 | H19400 | AB029000 | AB020713 |
| | Affy | 49599_at | 88138_at | 63393_at | 76253_at | 52793_at | 90920_at | 35832_at | 41812_s_at |
| | SedID | 1194 | 250 | 302 | 334 | 350 | 986 | 278 | 276 |
| | #1 | 254 | 255 | 256 | 257 | 258 | 259 | 260 | 261 |

| | | | | | 7 | 2 | | | |
|---|---------------------|---|---|--|---|---|--------------------------|---|--|
| | p-values | 9.6801E-06 | 0.04190137 | 2.62E-08 | 5.93E-09 | 5.5307E-05 | 3.56E-09 | 0.0000146 | 0.01054263 |
| | Fold Change | 0.247630859 | 0.33 | 0.226047115 | 0.075355998 | 0.22 | 6.816530863 | 0.325833287 | 0.3274967 |
| | Cluster Description | Cluster Ind. AA142956:z443g02.s1 Homo sapiens cDN4, 3 end /donne-IINA/GE-504722 idone_end=3' (gb=AA142956 /gi=1712334 /lug=ls_239142_Inen=640* | Cluster Ind. AA701619:zi41f11.s1 Homo sapiens cDN4, 3 end /clone=433389 /clone_end=3 /gb=AA701619 /gi=2704784 /ug=Hs.190382 Ine=398 | Cluster Incl AB011089:Homo sapiens mRNA for KIAA0517 protein, partial cds (cds=(0,2380) /gb-AB011089 /gl=3043557 /ug=Hs.12372 /en=5740 | Cluster Incl AB002351;Human mRNA for KIAA0353 gene, partial cds /cds=(0,4125) /gb=A8002351 /gi=2224646 /ug=Hs.10587 /elr=6551 | Hs.115740 Cluster Incl. AA706612:ah26c02.s1 Homo sapiens cDNA, a rend Idone=1238938 Idone_end=3'/gb=AA706612'/gi=2716530 Iug=Hs.115740 Jen=949 | | Cluster Incl D14664:Human mRNA for KIA40022 gene, complete cds (cds=(184,696) /dp=D14664 /dj=285952 /ug=Hs.2441 /len=3694 | Cluster Incl. Al881452:wj49b03.xt Homo sapiens cDNA, 3 end /clone=IMAGE-2406125 /clone_end=3' /gb=Al831452 /gj=5452123 /ug=Hs.91539 /len=597' |
| ĺ | Cluster# | Hs.42676 | Hs.7314 | Hs.12372 | Hs.10587 | Hs.115740 | Hs.81892 | Hs.2441 | Hs.111758 |
| | Gene Name | KIAA0781 protein | KIAA0614 protein | KIAA0517 protein | KIAA0353 protein | KIAA0210 gene product | KIAA0101 gene product | KIAA0022 gene product | keratin 6B |
| | Genbank | AA142956 | 68283_r_at AA701619 | AB011089 | AB002351 | AA706612 | D14657 | D14664 | Al831452 |
| | Affy | 65593_at | 68283_r_at | 39382_at | 39544_at | 91405_at | 38116_at | 34760_at | 62998_at |
| | SeqD | 28 | 203 | 273 | 269 | 207 | 961 | 962 | 731 |
| | #4 | 262 | 263 | 264 | 265 | 266 | 267 | 268 | 269 |

| WO 02/059 | 2/1 | | 7 | 3 | | PC 1/050 | 2/021/6 |
|---|------------------------|--|---|---|---|---|---------------------------------|
| p-values 0.00012792 | 0.0007796 | 7.6279E-07 | 1.23E-05 | 0.01689309 | 1.29E-07 | 0.0007917 | 6.2389E-06 |
| Fold Change 0.18 | 3.175184161 | 0.146596341 | 5.51416519 | 0.25 | 0.171820134 | 0.310324615 | 0.223459008 |
| Cluster Description Cluster Incl. Al566193':469902.x1 Homo septens cDNA, 3 end Iotone=IMAGE-2214075 Iotone_end=3' (pp=Al566193 /gj=4524645 Iuu-Hs, 198927 Ann=389 | | Cluster Incl. AW008322:wx53407.x1 Homo sapiens cDNA, 3 end /clone=IMAGE-890848 /clone_end=3 /gb=AW008322 /gi=5857100 /tu=he_234145 /len=556 | Cluster Incl AA203213::xx57e04.r1 Homo saplens cDNA, 5 end /clone=INAGE446622 /clone_end=6" /gb=A203213 /gi=1798923 /ug=Hs, 883 /len=879" | Cluster Incl. Al971748:wr07f08.x1 Homo sapiens cDNA, 3 end /clona=INAGE:2480871 /clona_end=3 /gb=Al971748 /gi=5768574 /uc=Hs_236704 /lorm=641* | Claster for AL021786; Human DNA sequence from PAC debetize on advancesome XQZ1.1- 21.2. Contains a mouse E26 like gene, a Kinesin like pseudogene and ESTs (cdse)(cg08) (pds-AL027786) (gj=2883186 Alg=lk; 7110) Alen=1389 | Cluster Incl. AW022660:df42a02.y1 Homo saplens cDNA, 5 end /clone=INA/GE-2485899 flone_end=6 /gb=AW022660 /gi=5876190 /ug=Hs.28002 /len=605 | |
| Cluster # Hs.196927 | Hs.237868 | Hs.64310 | Hs.833 | Hs.85266 | Hs.17109 | Hs.85112 | Hs.143648 |
| Gene Name iroquois homeobox protein 4 | interleukin 7 receptor | interleukin 11 receptor, alpha | interferon-stimulated protein, 15 kDa | integrin, beta 4 | irtegral membrane protein 2A | insulin-like growth factor 1 (somatomedia C) | insulin receptor substrate 2 |
| Genbank Al566193 | Al655781 | AW008322 | AA203213 | Al971748 | AL021786 | AW022660 | AA031286 |
| Affy 79663_at | 63628_at | 89921_at | 38432_at | 73285_i_at | 40775_at | 64305_s_at | 56338_at |
| SeqID 518 | 557 | 928 | 83 | 826 | 850 | 942 | 4 |
| 270 | 271 | 272 | 273 | 274 | 275 | 276 | 277 |

| | | 74 | | | |
|---|---|---|--|--|---------------------------------|
| p-values 0.0000161 | 0.0016656 | 0.000411 | 0.00034019 | 1.7517E-05 | 3.744E-08 |
| 6.309888618 | 0.316720069 | 0.23941203 | 0.302724058 | 10.30345974 1.7517E-05 | 0.1264716 |
| Cluster incl. ALOZZZSS-Human DNA sequence Cluster incl. ALOZZZSS-Human DNA sequence from clone 625H16 on chromosome 6922.2-23. Conclusis the gene for ID4 Helix-loop-helix DNA binding profiles and part of an alternatively spike-ad novel gene. Containe ESTs, 51'5s, 625s, and purely CQC islands (cise-1638, 853) (pp-ALOZZZZS (pj-8-6762777 /ug+1s, 3-4853 /Aer-3859 | Hs.204038 Cluster Inci. AA152276:206e06.r1 Homo sapiens cDNA, 5 end (donc=INAGE-491554 follow_2 and=5' (pg=A152276 fg=1721679 inc=+s.204038 lan=518 | Cluster Incl A1680656,wt23-07.x1 Homo cluster Incl A1680656,wt23-07.x1 Homo saplens cDN4.x and (Johne-IMAGE-2351436 Jchone_end=2" (ga-A1680656 /gi=4764239 Iug=Hs.78325 /len=552" | Hs.283305 Cluster Incl. Al813866:w/k61b03.x1 Homo saplens cDNA, 3 end /clone=IMAGE-2418863 /clone.gnd=2', Jope-M813866 /gi=5425081 /nr=He 2'A8013 /lon=530 | Cluster Incl. Al828404:wk84f07.x1 Homo sapiens cDNA, 3 end /clone=IMAGE-2422117 /clone endrel's (pg=5449075 /incle 2000x7 /incle | |
| Cluster # | Hs.204038 | Hs.76325 | Hs.283305 | Hs.140 | Hs.112844 |
| Gene Name inhibitor of DNA binding 4, dominant negative helix-loop- helix protein | indolethylamine N- methyltransferase | immunoglobulin J polypeptide, linker protein for immunoglobulin alpha and mu polypeptides | immunoglobulin heavy contant alpha 1 | immunoglobulin heavy constant gamma 3 (Gm | hypothetical protein PRO2160 |
| Genbank AL022726 | AA152276 | AI660656 | Al813866 | AI828404 | AW016235 |
| Aff <u>y</u> 41536_at | 63798_at | 37006_at | 75127_f_at | 58869 <u>i</u> at | 64258_f_at |
| SeqID 852 | 29 | 570 | 989 | 727 | 935 |
| 278 | 279 | 280 | 281 | 282 | 283 |

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|---------------------|---|------------------------------|--|----------------------------------|----------------------------------|----------------------------------|----------------------------------|----------------------------------|
| b-values | 2.1399E-07 | 4.8443E-05 | 1.7499E-09 | 5.2039E-08 | 0.00014333 | 0.00241384 | 1.8721E-05 | 0.00187809 |
| Fold Change | 0.297383187 | 3.983397486 | 0.047728145 | 0.289179561 | 3.336721416 0.00014333 | 0.32 | 3.188106939 | 4.36 |
| Cluster Description | Cluster Incl. AA651733:ns87b07.r1 Homo sapiens cDNA, 5 end /clone=ll/AAGE-1188661 /clone_end=67 (gb=A651733 /gi=2583385 /ug=Hs.21861 /Ren=515 | | | | | | | |
| Cluster# | Hs.21861 | Hs.42949 | Hs.271277 | Hs.169764 | Hs.118552 | Hs.267368 | Hs.133260 | Hs.263081 |
| Gene Name | hypothetical protein MPMGp800C04260Q 003 | hypothetical protein HES6 | hypothetical protein from EUROIMAGE 363668 | hypothetical protein FLJ20701 | hypothetical protein FLJ20539 | hypothetical protein FLJ20489 | hypothetical protein FLJ20354 | hypothetical protein FLJ20320 |
| Genbank | AA651733 | N21131 | AA020743 | AA521440 | AI276023 | AA563601 | AA813827 | A1149537 |
| Affy | 60810_at | 49633_at | 50223_at | 54742_at | 47427_at | 91394_at | 56634_at | 70129_r_at |
| SedID | 195 | 1063 | ത | 144 | 406 | 167 | 231 | 371 |
| # | 284 | 285 | 286 | 287 | 288 | 589 | 290 | 291 |
| | | | | | | | | |

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|---|----------------------------------|---|--|---|--|--|----------------------------------|----------------------------------|
| p-values 0.00010396 | 0.00090349 | 6.097E-08 | 3.4283E-07 | 1.2408E-05 | 0.00013093 | 9.6896E-05 | 1.4039E-06 | 0.00250076 |
| Fold Change 3.437531315 | 4.739571659 | 3.646875259 | 0.299454892 | 0.263536613 | 0.223334561 | 0.142325635 | 3.915587607 | 0.33 |
| Cluster Incl. AA181080:zp68h06 s1 Homo sapiens cDNA, 3 end (olone=IMAGE-825211 clone=end=3'(pb=A181080'gj=1764543 /ug=hs.239302,len=807. | | Cluster Incl. AA741298:oc86a07.s1 Homo saplens cDNA /clone=IMA/GE-1356564 /cgb-AA741298 /gi=2779890 /ug=Hs.23495 /len=530 | Cluster Incl. Al376944:tc34b06.x1 Homo sapiens cDNA, 3 end /clone=IMA/GE-2066483 /clone_end=3' /gb-A376944 /gi=4186797 /ucHs 61 /66 /len=482". | Cluster Incl. W27378:28b7 Homo sapiens cDNA /gb=W27376 /gj=1307036 /ug=Hs.8395 /len=596 | Cluster Incl. A1971695:wq88e01.x1 Homo saplens CDNA, 3 and /clone=INAGE:-2479128 /clone_end=3'/gb=A1971695/gi=5768521 /ug=Hs.237607/len=573° | Cluster Incl. AA708740:z157g12.s1 Homo sapiens cDNA, 3 and /clone=508086 /clone_end=3 /gb=AA708740 /gi=2718658 /uc=Hs.23467 / Herr=475 | | |
| Cluster# Hs.92254 | Hs.161554 | Hs.23495 | Hs.6166 | Hs.8395 | Hs.107882 | Hs.23467 | Hs.26516 | Hs.128766 |
| Gene Name hypothetical protein FLJ20163 | hypothetical protein FLJ20159 | hypothetical protein FLJ11252 | hypothetical protein FLJ11196 | hypothetical protein FLJ10781 | hypothetical protein FLJ10659 | hypothetical protein FLJ10633 | hypothetical protein FLJ10604 | hypothetical protein FLJ10600 |
| Genbank AA181060 | AI766666 | AA741298 | Al376944 | W27376 | Al971695 | AA708740 | AA195220 | AI811688 |
| Affy 50280_at | 74539_g_at | 43439_at | 53962_at | 55074_at | 44855_s_at | 44057_at | 51130_at | 74343_at |
| SeqID 76 | 656 | 213 | 445 | 1179 | 825 | 210 | 79 | 989 |
| ## 292 | 293 | 294 | 295 | 296 | 297 | 298 | 599 | 300 |

| | | | | | 7 | 7 | | | |
|---|---------------------|---|---|----------------------------------|----------------------------------|----------------------------------|----------------------------------|---|----------------------------------|
| | p-values | 0.0022939 | 4.719E-12 | 1.3006E-07 | 9.0609E-08 | 1.8056E-05 | 7.6457E-06 | 0.00028995 | 5.3484E-07 |
| | Fold Change | 3.07 | 9.164231149 | 3.058813358 | 4.806177129 | 3.509937557 | 0.21534971 | 3.782457573 | 0.18625739 |
| | Cluster Description | Cluster Incl. Al016073:ov26ff1.xf Homo sepiens ColbN, 3 end /clone=INAGE-1638477 /iclone_end=3' /gb=Al016073 /gj=3230409 /lug=Hs.131640 /len=561* | Cluster Incl. Al674163:wc08a11.x1 Homo sapiers cDNA, 3 end /clone=INAGE-2314662 /clone_end=3' /gb=Al674163 /gi=4874643 /ug=Hs.14559 /len=553' | | | | | Cluster Incl. Al949698:wq13h04.x1 Homo sapiens cDNA, 3 end /clone=INA/GE-247191 /clone_end=3 /gb=Al949698 /gl=5742008 /uc=Hs.42309 /eln=653 | |
| , | Cluster# | Hs.131840 | Hs.14559 | Hs.48855 | Hs.122579 | Hs.104650 | Hs.106283 | Hs.53913 | Hs.22505 |
| | Gene Name | hypothetical protein FLJ10578 | hypothetical protein FLJ10540 | hypothetical protein FLJ10468 | hypothetical protein FLJ10461 | hypothetical profeln FLJ10292 | hypothetical protein FLJ10262 | hypothetical protein FLJ10252 | hypothetical protein FLJ10159 |
| | Genbank | Al016073 | AI674163 | AA134589 | AI823992 | Al037879 | Al651535 | Al949698 | AL040063 |
| | Affy | 74690_at | 58235_at | 52966_at | 59461_at | 54889_at | 64666_at | 52617_at | 49523_at |
| | SeqID | 313 | 582 | 53 | 715 | 323 | 549 | 801 | 863 |
| | #1 | 301 | 302 | 303 | 304 | 305 | 306 | 307 | 308 |
| | | | | | | | | | |

| | WO 02/0592 | 271 | | 7 | 8 | | PCT/U | S02/02176 |
|---------------------|--|--|---|---|---|--|---|--|
| p-values | 5.8315E-11 | 0.00017758 | 0.00279468 | 1.5285E-06 | 5.019E-05 | 5.4416E-07 | 0.00026772 | 1.2153E-08 |
| Fold Change | 4.802394237 | 0.27 | 0.26 | 0,262634843 1,5285E-06 | 0,269743016 | 0.275006365 | 0.291337727 | 0.054106026 |
| Cluster Description | Cluster Ind. AA779101;2/42trl2.s1 Homo sapiens ChONA, 3 et d/done=452999 folone_end=3' (gb=AA779101 /gl=2838432 /lgf=16:104695) flan=528 | Cluster Incl. Al613455:ty37b03.x1 Homo sspiens pDNA, 3 end /done=IMAGE-2281229 /clone_end=3' /gb~Al613455 /gj=4622622 /tg=18: 181658 len=418' | Cluster Incl. At394248:tf78b06.x1 Homo sapiens CDNA, 3 end /done=IMAGE-2105383 /done_end=3' /gb=Al39428 /gi=4223795 //ge18.235588 Men=590' | Cluster Incl. Z78379:HSZ78379 Homo sapiens cDNA /clons=3.18-(CEPH) /gb=Z78379 /gi=1495152 /ug=Hs.224337 /len=1958 | Cluster Inci. AL039400:DKFZQ434K1210_s1 Homo sapiens cDNA, 3 end Homo sapiens cDNA, 3 end Home_BNFZQ434K1210 /done_end=3' fgb=AL039400 /gj=S928564 /ug=Hs.32352 flen=703' | Cluster Ind. Al632223:tt20d07.x1 Homo sepiens cDN4, 3 enf clone=IMAGE-2241325 fclone_end=3'/gb=Al632223/gi=4683553 fgi=4684633 lug=16; 44633 llen=721' | Cluster Ind. AA625897.zu87b02.s1 Homo sapiens cDNA, 3 end /done=IMAGE-744939 idone_end=3' /gb=AA625897 /gj=2538284 /ug=16, 222095 /len=252' | Cluster Incl. AA048671:zf12d09.r1 Homo sepiens cDNA, 5 end /clone=IMAGE-376721 /clone_end=5 /gb=AA048671 /gi=1524772 /ug=Hs.17917 /len=596' |
| Cluster # | Hs.104859 | Hs.49933 | Hs.284280 | Hs.279023 | Hs.32352 | Hs.144633 | Hs.24583 | Hs.278975 |
| Gene Name | hypothetical protein DKFZp762E1312 | hypothetical protein DKFZp762D1011 | hypothetical protein DKFZp547H236 | hypothetical protein DKFZp434N1928 | hypothetical protein DKFZp434K1210 | hypothetical protein DKFZp434F2322 | hypothetical protein DKFZp434C0328 | hyaluronic acid receptor,lymphatic vessel endothelial hyaluronan receptor 1 |
| Genbank | AA779101 | Al613455 | Al394248 | Z78379 | AL039400 | AI632223 | AA625897 | AA046671 |
| Affy | 48045_at | 77581_at | 87842_at | 63581_at | 44682_at | 53831_at | 72538_at | 65731_at |
| SedID | 225 | 533 | 460 | 1245 | 859 | 939 | 186 | 56 |
| #1 | 308 | 310 | 311 | 312 | 313 | 314 | 315 | 316 |

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|---------------------|---|-----------------|---|---|---|---|--|---------------------------------|
| b-values | 2.013E-09 | 0.00165492 | 0.00126636 | 0.00508853 | 6.2245E-06 | 1.93E-08 | 2.4107E-05 | 0.0031179 |
| Fold Change | 7.363245733 | 0.26 | 3.975570707 | 3.439858165 | 6.704063952 | 3.965279191 | 0.285742894 | 5.05 |
| Cluster Description | Cluster Incl. Al990409:w774803.x1 Homo sapiens cDN4, 3 end /done=IMAGE-2513164 clone_end=3'/gb=Al990409 /gj=5837290 /tig=ls,5199 /len=744* | | Cluster Incl. Al670876:wa06c12.x1 Homo saplens cDN4, 3 end /clone=IMAGE-2297302 /clone_end=3'/gb-48700876 /gj=4850607 /ug=Hs.44276 /ne=798' | Cluster Ind. AW024474:wu76h03.x1 Homo sapiens cDN4, 3 end /clone=IMAGE-990888 Tclone_end=3'/gb=Aw024474/gj=5878004 /ug=ls_236884 len=229' | Cluster Ind. AI525822:PT1.3_06_B10.r Homo sapiens cDNA, 5 end /clone_end=67 /gb=AI525822 /gj=4439957 /ug=Hs.109706 /fen=773 | Cluster Ind AA255502:zr85b06.r1 Homo sapiens cDNA, 5 end Iclone=INAGE-882451 Iclone_end=5" igb=AA255502 igi=1892406 fug=Hs.46423 inen=348" | Cluster Incl. AA609509:aff14d11.s1 Homo sapiens cDNA, 3 end /clone=IN/AGE-1031637 /clone_end=3 /gb=AA609509 /gi=2457937 /uc=Hs.28503 /len=381* | |
| Cluster# | Hs.5199 | Hs.278943 | Hs.44276 | Hs.44276 | Hs.109706 | Hs.46423 | Hs.180877 | Hs.180779 |
| Gene Name | HSPC150 protein similar to ubiquitin-conjugating enzyme | HSPC047 protein | homeo box C10 | hотео box C10 | HN1 protein | H4 histone family, member G | H3 histone, family 3B (H3.3B) | H2B histone family, member B |
| Genbank | Al990409 | F37133 | AI670876 | 57830_s_at AW024474 | AI525822 | AA255502 | AA609509 | AI125923 |
| Affy | 45803_at | 78942_at | 52117_at | 67830_s_at | 56428_at | 39969_at | 90048_at | 88155_at |
| SeqD | 842 | 27.5 | 573 | 947 | 503 | 91 | 179 | 356 |
| #1 | 317 | 318 | 319 | 320 | 321 | 322 | 323 | 324 |
| | | | | | | | | |

| | | | | | 80 | | | | |
|---------------------|---|---|--|--|---|--|---|----------------------------------|---------------|
| p-values | 5 | 1.5236E-07 | 3.9853E-10 | 2.305E-06 | 8,5076E-08 | 3.05 E- 08 | 2.6578E-09 | 1.3623E-06 | 1.8509E-07 |
| Fold Change | | 0.226916444 | 4.119810176 | 0.172426013 | 12.40 | 0.259661725 | 0.233804467 | 0.17 | 5.46370987 |
| Cluster Description | Cubisis Into N.3 end /done=INAGE-2429487 septems CDNA, 3 end /done=INAGE-2429487 /done end-3"/gb=/R85852 /gj=5591016 /dg=Hs.795 /len=580" | Cluster Ind. A4115300;z/08d12.s.1 Homo sapiens cDNA, 3 end /done=IMAGE-501431 /done_end=3' /gb=A4115300 /gj=1670497 /lg=18,103720 /len=588 | Cluster Ind. U82984:U82984 Homo sapiens cDNA /clone=163g24 /gb=U82984 /gi=2731436 /ug=Hs.23900 /len=1771 | Cluster Ind. D60584:HUM118E12A Homo sapiens cDNV, 3 end /donne-GEN-118E12 /done_end=3'/gb=D60584/gi=962223 /ug=ls_234684/len=341' | Cluster Incl. AA613715:nq25b01.s1 Homo sapiens cDNA, 3 end /donne-IMAGE-1144873 /clone_end=3' (gb=AA613715 /gj=2463685 /ug=18,81795 /flen=6077 | | Cluster Incl. AA524029.ng32f02.s1 Homo sapiens cDNA, 3 end /done=IMAGE-838507 /done_end=3 /gb=#A524029 /gi=2264957 /ug=Hs.77898 /len=721* | | |
| Cluster # | 69.3 | Hs.23767 | Hs.23900 | Hs.58589 | Hs.5566 | Hs.160318 | Hs.77889 | Hs.239069 | Hs.118162 |
| Gene Name | nzk filstone laffilly, member O | guanine nucleotide binding protein (G protein), gamma 2 | GTPase activating protein | glycogenin 2 | gap junction protein, beta 2, 26kD (connexin 26) | FXYD domain- containing ion transport regulator 1 (phosoholemman) | Friedreich ataxia region gene X123 | four and a half LIM domains 1 | fibronectin 1 |
| Genbank | Alososoz | AA115300 | U82984 | D60584 | AA613715 | AA524547 | AA524029 | AW024276 | AW021977 |
| Affy | 32609_at | 58445_at | 50271_at | 55630 <u>f</u> at | 91306_s_at | 32109_at | 54581_at | 80572_at | 45557_r_at |
| SeqD | 96/ | 40 | 1163 | 696 | 182 | 149 | 145 | 944 | 940 |
| ## G | 975 275 | 326 | 327 | 328 | 329 | 330 | 331 | 332 | 333 |

| | | | | 8 | 1 | | | |
|---------------------|--|---|--|--|---|---|--------------------------|--------------------------|
| p-values | 5.0303E-07 | 2.9946E-05 | 0.00020319 | 0.00010017 | 6.067E-07 | 0.00101117 | 6.263E-08 | 7.8076E-06 |
| Fold Change | 0.097550234 | 3.294370164 | 0.276576877 | 0.31 | 0.08111725 | 0.311274497 | 0.108158864 | 0.22460642 |
| Cluster Description | Cluster Inci. AA156998:z119e05.s1 Homo sepiens cDNA, 3 end /olone=IMAGE-502400 /olone_end=2/ igb=AA156998 /gj=1728613 /ug=Hs.239679 /len=562' | | Cluster Incl. AA224344:zr18d12.sr1 Homo sapiens cDNA, 3 end (clone=IMAGE-683575 febre_end=3' (gb=AA224344 /gj=1844967 /ug=18.237937 Nen-420' | Cluster Incl. A1138998:qd83f09.x1 Homo sapiens cDNA, 3 end (clone=INAGE-1738105 fclone_end=3' (gb=A113898 /gl=3644970 fug=16.234468 len=570° | Cluster Incl. A1912678:we12e04.x1 Homo saplens cDNA, 3 end /clone=IN/AGE-2340894 /clone_end=3' /gb=A1912678 /gl=5632533 /tu=Hs.11713 /fan=587 | Cluster Incl. AA557237.n175d07.s1 Homo saplens cDNA, 3 end /clone=INAGE-1056493 Aclone_end=3 /gb=A4557237 /gi=2327714 /ug=Hs, 168776 /len=605 | | |
| Cluster# | Hs.211568 | Hs.193053 | Hs.43697 | Hs.79095 | Hs.11713 | Hs.21595 | Hs.109439 | Hs.108924 |
| Gene Name | eukaryotic translation initiation factor 4 gamma, 1 | eukaryotic translation initiation factor 2C, 2 | ets variant gene 5 (ets-related molecule) | epidermal growth factor receptor pathway substrate 15 | E74-like factor 5 (ets domain transcription factor) | DNA segment on chromosome X and Y (unique) 155 expressed seguence | DKFZP586P2421 protein | DKFZP586P1422 protein |
| Genbank | AA156998 | A1146465 | AA224344 | A1138998 | Al912678 | AA557237 | AW020116 | W72194 |
| Affx | 63893_f_at | 56226_st | 75258_f_at AA224344 | 78641_at | 59390_at | 60058_at | 43506_at | 56409_at |
| SealD | 02 | 369 | 87 | 363 | 768 | 166 | 937 | 1197 |
| # | 334 | . 332 | 336 | 337 | 338 | 339 | 340 | 2 . |

| | WO 02/0592 | 271 | | 8 | 2 | | PCT/US0 | 2/02176 |
|---------------------|--|--|--|---|---|--|---------------|----------------------------------|
| p-values | 7.5151E-06 | 6.4105E-05 | 0.00232454 | 0.00000163 | 6.3581E-06 | 0.00455471 | 0.000426 | 4.57E-06 |
| Fold Change | 0.185480277 | 0.308559272 | 0.306209896 | 0.296976926 | 3.405694621 | 0.222139001 | 0.215553985 | 5.17065685 |
| Cluster Description | Cluster Ind. W78050.zd78c04.s1 Homo saplens CDNA, 3 end /clone=IMAGE-346758 /clone_end=3' /gb=W78050 /gl=1388613 /ug=He,236316' Nen-446' | Cluster Incl. AW026659:wv15c11.x1 Homo saplens cDNA, 3 end Icloine=IMAGE-991047 Icloine_end=3' (gb=AW026659 /gj=5880112 Mg=18_29358 /fen=591' | Cluster Incl. A1984087:wz56d08.x1 Homo sapiens cDN4, 3 erd /clone=IMAGE-2562083 /clone_end=3 /gb=A1984087 /gi=5811306 /ug=He.235102.llen=479 | Cluster Incl AL050024:Homo sapiens mRNA; cDNA DKF2p5840206 (from done DKF2p5840206) cds=(0,404) /gp=AL050024 /gp=A88A093 Aug=Hs 25956 inen=1409 | Cluster Incl. AW007289:wt54f03.x1 Homo saplens cDNA, 3 end /dona=IMAGE-2511293 /done_end=3' /gb=AW007289 /gj=5855067 /ug=Hs. /6441 / len=552 | Cluster Incl. A1961431:wt22e08.x1 Homo saplens ColbA, 3 end /clone=IMAGE-2508230 /clone_end=3' /gb=A1961431 /gl=5754144 /ug=Hs.9029 /en=696' | | |
| Cluster# | Hs.58419 | Hs.26358 | Hs.3447 | Hs.25956 | Hs.16441 | Hs.9029 | Hs.80552 | Hs.184641 |
| Gene Name | DKFZP586L2024 protein | DKFZP566K1924 protein | DKFZP564K1964 protein | DKFZP564D206 protein | DKFZP434H204 protein | DKFZP434G032 protein | dermatopontin | delta-6 fatty acid desaturase |
| Genbank | W78050 | AW026659 | A!984087 | AL050024 | AW007289 | Al961431 | AL049798 | AL050118 |
| Affy | 44813_s_at | 64180_at | 45501_s_at | 39577_at | 60038_at | 48684_at | 38057_at | 32190_at |
| SeqID | 1208 | 951 | 839 | 198 | 922 | 128 | 988 | 892 |
| 啪 | 342 | 343 | 344 | 345 | 346 | 347 | 348 | 349 |

| | | | | : | 83 | | |
|-------------|---|---|---|--|--|--|---|
| p-values | 1.3941E-05 · | 0.00379696 | 0.00136947 | 1.95-08 | 0.00679779 | 0.00061484 | 5.3744E-06 |
| Fold Change | 0.23 | 4.29 | 0.24 | 0.121682021 | 3.01095294 | 3.125264866 | 0.297772176 |
| | Cluster Ind. A4459140:tj65e06.x1 Homo sapiens cDNA, 3 end /done=IMAGE-2146402 /done_end=3' /gb=A459140 /gi=4311719 /lg=16:129109 /len=499 | Cluster Ind. AA909181:ol12b04.s1 Homo sapiens cDN4, 3 end /olone=IMAGE-1522215 /clone_end=3'/gp=AA909181 /gi=3048586 /ug=He,234830 /len=528 | Cluster Ind. X84721.HSEST222 Homo sapiens cDNA /done=MEC-222 /gb=X84721 /gi=673398 /ug=Hs.164886 /len=558 | Cluster Incl AL038340.DKFZp596K192_s1 Homo saplens cDN4, 3 end (Homo=DKFZp568K192 /done_end=3" (gb=AL038340 /gl=5407591 /ug=Hs.1940 Nen=748" | Cluster Incl. AA127736:zk88c12.r1 Homo saptens cDNA, 5 end /clone=IMAGE-488910 /clone_end=5/ /gb=AA127736 /gj=1687099 /ug=Hs,237523 Hen=61 f9 | | Classir Inc. (2824415:874204.x1 Homo sepiens cDNA, 3 end domenlAAGE-2049683 dome_end=2' /gb=40882415 /gl=4195196 /ug=Hs_239510 /len=418 |
| Cluster# | Hs.74649 | Hs.1174 | Hs.164866 | Hs.1940 | Hs.82985 | Hs.179573 | Hs.75106 |
| Gene Name | cytochrome c oxidase subunit Vic | cyclin-dependent kinase inhibitor 2A (melanoma, p16, inhibits CDK4) | cyalin K | crystallin, alpha B | collagen, type V, alpha 2 | collagen, type I, alpha Hs.179573 2 | clusterin (complement Hs.75106 19.8is Intibiot, SP-40,40, sulfated glycoprolein 2, glycoprolein 2, existeriorne- repressed prostate message 2, apolipoprotein J) |
| Genbank | Al459140 | AA909181 | X84721 | NM_001885 | AA127736 | AI610692 | Al382416 |
| Affy | 74406_at | 73132_r_at AA909181 | 90629_at | 32242_at | 65797_at | 49162_f_at | 75384_ <u>f_at</u> |
| SedID | 481 | 247 | 1236 | 1100 | 4 | 531 | 457 |
| 妣 | 350 | 351 | 352 | 353 | 354 | 355 | 326 |

| | | | | 84 | | | |
|------------------|---------------------|---|--|---------------------|--|---|---|
| | p-values | 2.3 5 79E-06 | N | 0.000347 | 1.1789E-10 | 0.00000214 | 2.8896E-05 |
| | Fold Change | 0.262336411 | 4.805037919 | 0.212939754 | 8.862541971 1.1789E-10 | 0.155185151 | 3.067793027 |
| m r | Cluster Description | Clanse Ind. (18824 f. Ist.)2004 xt. Homo sapiers CDNA, 3 end. (drone=INAOE-204688) fobre, end-3' (gb=A1882415 (gi=4185196 /ug=Hs_289510 /len=418* | Hs.279905 Cluster Incl. AA143745:zo31a01.s1 Homo sapiens cIDNA, 3 end /dons=INIAGE-888456 /done_end=3/gb=AA143745/gi=1713158 /ug=Hs.62273 len=B49 | | Cluster Ind. AA534688:nf75c01.s1 Homo septens cDNA, 3 end folone=INA/GE-925728 Idone_end=3' /gib-A534688 /gi=2278941 /ug=1e,238549 hen-467" | Cluster Ind ALD49176:Human DNA sequence from dove 14/16 on dinomasone AC21-4-2. Gondains parks of a novel Chordin LIKE protein with von Willebrand factor type C domains. Contains ESTs, STSs and GSSs lods-40/167), dip—ALD49176 (gi—4808228 /ug=Hs, 82223 /ug=Hs, 82423). | Cluster Incl. AA426499:zw02b06.r1 Homo sapiens cDNA, 5 end /clone=IMAGE-768083 /done_end=5'/gb=AA426489 /gi=2106744 /ug=Hs.239900 /len=553 |
| The Committee of | | Hs.75106 | Hs.279905 | Hs.162209 | Hs.9329 | Hs.82223 | Hs.81800 |
| | Gene Name | clusterin (complement lysis inhibitor, SP- 40,40, sulfated glycoprotein 2, testosterone- repressed prostate message 2, politoproriein J) | clone HQ0310 PRO0310p1 | claudin 8 | chromosome 20 open reading frame 1 | chordin-like | chondroitin sulfate proteoglycan 2 (versican) |
| | Genbank | Al382415 | AA143745 | AL049977 | AA534688 | AL049176 | AA426499 |
| | Affy | 75382 <u>i_</u> at | 64489_at | 33611_g_at AL049977 | 45574_g_at AA534688 | 37630_at | 45718_at |
| | SedID | 457 | 09 | 888 | 158 | 882 | 13 |
| | # | 367 | 358 | 326 | 360 | 361 | 362 |
| | | | | | | | |

| | | | 85 | | | | |
|--|---|---|---|---|--|--|--|
| p-values 7.4702E-06 | 8.7397E-08 | 1.4906E-07 | 7.4051E-05 | 0.03811473 | 1.16 E-0 7 | 0.00000833 | 3.194E-05 |
| Fold Change 3.05 | 0.174212976 | 5.21 | 0.221161273 | 3.355023106 | 0.231001071 | 0.292109229 | 0.23945116 |
| Cluster # | Cluster Ind. AA545730:HBMSF2G12-REV Homo sapiens cDNA, 5 end cloine-HBMSF2G12, Idone, end=5 (gb-AA545730) (gi=2307100 /ug=Hs.31198 | | Cluster Ind. Al972237:wr33c02.x1 Homo sepiens cDN4, 3 end /clone=IMAGE-2489474 fictore_end=3 /gb-Al972237 /gl=5769063 /tuc+ls_233663 flen=354* | Cluster Ind. A675178:tm80g06.x1 Homo sapiens cDN4, 3 end Iclone=IMAGE-2164774 Iclone=end=3'/gb-A675178 /gj=4875658 Iuc+15, 50207 Ihen=462 | Cluster Incl AB020629:Homo sapiens mRNA for NIAA0822 protein, complete cds /cds=(138,4883) /gb=AB020629 /gi=4240129 /tic=ls_38095 /len=5677 | Cluster Incl AI651024;wa96h06.x1 Homo sapiens cDNA, 3 end Idone=IMAGE-2304059 Idone-end=3" Igb=AI651024 Igi=4735003 Iuc=Hs, 15780 Nen=657" | |
| Cluster# Hs.240443 | Hs.117582 | Hs.184572 | Hs.82129 | Hs.90207 | Hs.38095 | Hs.15780 | Hs.153961 |
| Gene Name chondrollin 4- sulfotransferase,chon drollin-4- sulfotransferase | CGI-43 protein | cell division cycle 2, G1 to S and G2 to M | carbonic anhydrase III, muscle specific | calclum channel, voltage-dependent, gamma subunit 4 | ATP-binding cassette, sub-family A (ABC1), member 8 | ATP-binding cassette, sub-family A (ABC1), member 6 | ARP1 (actin-related protein 1, yeast) homolog A (centractin alpha) |
| Genbank AA766775 | AA545730 | AF154332 | AI972237 | Al675178 | AB020629 | Al651024 | 64423_s_at AA628405 |
| Affy 75722_at | 50177_at | 91194_at | 74571_s_at Al972237 | 62987_r_at | 35717_at | 35390_at | 64423_s_at |
| SeqID 217 | 164 | 310 | 828 | 586 | 275 | 547 | 188 |
| 3€3 | 364 | 365 | 366 | 367 | 368 | 369 | 370 |

| | | WO 02/059 | 271 | | 8 | 66 | | PC | 1/US02/02176 |
|---|---------------------|---|--|----------------------------------|---|--|---|--------------------------------|---|
| | b-values | 3.3556E-06 | 1.4825E-06 | 2.0018E-05 | 0.000333 | 0.0000909 | 2.9261 E -05 | 0.0000126 | 0.00059716 |
| | Fold Change | 0.212140139 | 3.58350616 | 0.263359832 | 0.215928239 | 0.129092155 | 0.135508105 | 0.308528713 | 0.33 |
| ! | Cluster Description | Cluster Incl. Al804914.tu43d04.x1 Homo sapiens cDNA, 3 end /done=IMAGE-2253799 fidone_end=3'/gp-Al804914 /gj=5391504 /ug=18,55665 /len=356" | Cluster Incl. Al341261:qx85e07.x1 Homo sepiens cDNA, 3 end /clone=IMAGE-2009268 /clone_end=3'/gb=Al341261 /gi=4078188 /tig=1s,62180 /Ren=538' | | Cluster Incl Al381790:te41h10.x1 Homo saplens ClothA, 3 end /done=IMAGE-2089315 /clone_end=3" /gb=Al381790 /gl=4194571 /(ig=15.74120 /gl=544" | Cluster Incl H15814;y/28b07.s1 Homo sapiens cDNA, 3 and /clone=IMAGE-159541 /clone_end=3" /gb=H15814 /gl=880634 /ug=H5.90495 /len=453" | Cluster Incl. AA393277-zf74d08.r1 Homo sapiens cDNA, 5 and /clone=IMAGE-728079 fclone_and=5 /gb=AA393277 /gj=2046245 /uc=15.238152 han-455 | | Cluster Incl. F37480.HSPD38277 Homo saplens cDNA Idone-sH-000003-0/096 igb=F37480 igi=823106 kig=Hs.221714 Ien=408 |
| i | Cluster# | Hs.55565 | Hs.62180 | Hs.203299 | Hs.74120 | Hs.80485 | Hs.18268 | Hs.158203 | Hs.58324 |
| | Gene Name | ankyrin repeat domain 3 | anillin | amylase, alpha 2A; pancreatic | adipose specific 2 | adipose most abundant gene transcript 1 | adenylate kinase 5 | actin binding LIM protein 1 | a disintegrin-like and metalloprotease (reprolysin type) with thrombospondin type 1 motif, 5 (aggrecanase-2) |
| | Genbank | Al804914 | Al341261 | Al218026 | Al381790 | H15814 | AA393277 | D31883 | F37480 |
| | Affy | 46108_at | 46194_at | 89031_at | 32527_at | 40657_r_at | 58927_at | 40155_at | 80160_at |
| | SeqID | 685 | 426 | 392 | 455 | 982 | 101 | 965 | 978 |
| | 411 | 371 | 372 | 373 | 374 | 375 | 376 | 377 | 378 |

| wo | 02/ | 059 | 271 |
|----|-----|-----|-----|

| WO 02/059 | 271 | | | 87 | | PCT/ | US02/02176 |
|---|--|-------------|--|---|-------------|---------------------|------------|
| <u>p-values</u> 0.00028766 | 7.756E-06 | 0.00049057 | 1.5774E-06 | 2.3135E-06 | 3.4955E-06 | 2.2368E-07 | 3.8401E-05 |
| <u>Fold Change</u> 3.46460752 | 0.14023914 | 0.288607766 | 0.310758648 1.5774E-06 | 4.823648195 | 0.241006046 | 0.156366951 | 6.80 |
| Cluster Incl. AA135525-x109e04.s1 Homo Cluster Incl. AA135525-x109e04.s1 Homo Belbers CDMA, and Idonne-IMAGE-501438 Idone_end=3"(pl=4135525 (gl=1696573 Iug=Hs.55009 Iner=608 | Cluster Incl. Al088609:qb14e04.x1 Homo saplens cDNA, 3 end (clone=INAGE-1686254 /clone_end=3' (gb=Al088609 /gi=3427668 /uc=Hs.98556 /len=749 | | Cluster Incl. W37770:zc12g12.r1 Homo sapiens cDNA, 5 end /clone=IMAGE:322150 /clone_end=6 /gb=W37770 /gl=1319383 /ucHs, 9851 len=573* | Cluster Incl. AA14784:z150b04.s1 Homo sapiens cDNA, 3 end (clone=I)MAGE-505327 (clone_end=3 / gb=A4147884 /gi=1717300 /ureHs, 8812 /len=552 | | | |
| Cluster# Hs.56009 | Hs.98558 | Hs.98518 | Hs.9851 | Hs.9812 | Hs.97876 | Hs.97876 | Hs.97837 |
| Gene Name 2'-5'oligoadenylate synthetase 3 | | | | | | | |
| Genbank AA135525 | AI088609 | AW007983 | W37770 | AA147884 | Al970823 | AW016780 | Al935915 |
| Affy 64450_at | 58361_at | 75011_at | 55720_at | 49052_at | 63041_i_at | 45353_s_at AW016780 | 85521_at |
| SeqID 54 | 344 | 927 | 1184 | 63 | 820 | 936 | 792 |
| 379 | 380 | 381 | 382 | 383 | 384 | 385 | 386 |

| WO 02/059 | 271 | | 8 | В | | PC | 1/US02/0217 |
|---|-------------|---|--|------------------------|-------------|--|--|
| p-values 1.4682E-05 | 5.524E-07 | 8.975E-06 | 4.8195E-06 | 0.00058791 | 0.00034656 | 0.00068973 | 2.6139E-08 |
| <u>Fold Change</u> 0.161954139 | 0.281787455 | 0.302684307 | 0.12598837 | 0.248393008 0.00058791 | 3.039821602 | 3.308493975 0.00068973 | 0.282705156 |
| Cluster Incl. Al659076:t197408.x1 Homo sapplens cDNA, 3 and Idone=IMAGE-2249487 Idone_end=3'' (pb~8165976 ig=4'762646 Idone_end=2'' (pb~8165976 ig=4'762646 | | Cluster Incl. D55886:HUM405B01B Homo sapiens cDNA, 5 end (done=GEN-405B01 done=enE-1/gb-D55886 (gi=970283 hu=Hs, 5972 len=553 | Cluster Incl. R54660;yj74b11.s1 Homo sapiens cDNA, 3 end clonne=IMAGE-154461 clone_end=3 /gb=R54660/gi=819118 /ug=Hs,95511 /hg=427 | | | Cluster Incl. Al669212:wc13cr10.x1 Homo saplens CDNA, 3 and cloine=INAGE-2315058 Iclone_end=3' (gb=Al69212 /gi=4833986 /ug=Hs,29127 /len=596 | Cluster Incl. Al972873:wr44f08.x1 Homo saplens cDNA, 3 end /done=INA/GE-2490587 /clone_end=3 /gb=Al972873 /gi=5769699 /ug=Hs.9167 /len=594 |
| Cluster# Hs.97031 | Hs.96996 | Hs.9572 | Hs.95511 | Hs.94789 | Hs.92679 | Hs.92127 | Hs.9167 |
| Gene Name | | • | | | | | |
| Genbank Al659076 | Al672356 | D55886 | R54660 | Al743671 | AA292431 | Al669212 | Al972873 |
| Affy 51785_s_at | 63035_at | 47579_at | 55484_r_at | 47566_at | 46737_s_at | 55436_at | 65976_g_at Al972873 |
| SeqID 564 | 277 | 896 | 1112 | 635 | 83 | 572 | 832 |
| 387 | 388 | 389 | 390 | 391 | 392 | 383 | 394 |

| | | | | | | , | | | |
|--------------|---------------------|--|-------------|------------|-------------|--|--|--|-------------|
| | p-values | 3.1056E-07 | 0.00185863 | 0.00208182 | 0.00000576 | 5.7165 E-0 7 | 0.00000349 | 2.6349E-05 | 0.00010051 |
| | Fold Change | 0.146420991 | 3.433762656 | 0.29336996 | 0.290791559 | 0.159849377 | 0.316621071 | 0,20131865 | 0.306142564 |
| 1 - American | Cluster Description | Cluster Incl. Al972873:wr44f08.x1 Homo sapiens ChUN, 3 end /clone=INAGE-2490567 /clone_end=3'/gb=Al972873 /gi=5769699 /ug=Hs.9f67 /lene=594' | | | | Cluster Incl. AA351076:EST58700 Homo saplens ColbA, 3 end /done=ATCC-104314 ficine_end=3' /gb=AA351076 /gi=2003416 /ugHs.237155 /len=529 | Cluster Incl AL079279:Homo sapiens mRNA full length insert cDNA clone EUROIMAGE 248114 (cds=UNKNOWN /gb=AL079279 /gj=6102865 /ug=Hs.8963 ilen=2428 | Cluster Incl. AW006898:ws15g04.x1 Homo sapiens cDNA, 3 end /clone=INAGE-2497302 /clone_end=3' /pb=AW006898 /gi=5855676 /ug=Hs.224004 fen=228 | |
| | Cluster # | Hs.9167 | Hs.91109 | Hs.90756 | Hs.90063 | Hs.90063 | Hs.8963 | Hs.88827 | Hs.88537 |
| | Gene Name | | | | | | | | |
| | Genbank | AI972873 | AI742239 | Al150491 | AF052142 | AA351076 | AL079279 | AW006898 | AA481493 |
| | Affy | 6 597 5_at | 55610_at | 77001_at | 38803_at | 52294_s_at | 38786_at | 63994 <u> </u> at | 64813_at |
| | SedID | 832 | 629 | 374 | 297 | 88 | 894 | 919 | 132 |
| | #1 | 395 | 396 | 397 | 398 | 388 | 400 | 401 | 405 |
| | | | | | | | | | |

| | | | | 9 | 0 | | | -, |
|---------------------|--|-------------|-------------|---|-------------|------------|-------------|-------------|
| p-values | 1.3801E-05 | 6.8126E-06 | 1.0223E-07 | 2.2017E-05 | 4.7993E-06 | 2.8399E-09 | 5.6167E-06 | 1.8002E-06 |
| Fold Change | 0.224326303 | 0.319186626 | 4.319893329 | 0.300430737 | 0.233032608 | 0.16 | 0.113945577 | 0.204459738 |
| Cluster Description | Cluster Incl. AW007080:ws49h10.x1 Homo sapiens cDNA, 3 end Iclone=INA/GE-2500579 Iclone_end=3' Igb=AW007080 /gi=\$855858 Iug=Hs, 8817 Ilen=523 | | | Cluster Incl. AA480075:zv42b06.r1 Homo sapiens cDNA, 5 end (stone=II/MAGE-786275 /clone_end=5 /gb=AA480075 /gi=2208226 /ug=Hs.86016 flen=588* | | | | |
| Cluster# | Hs.8817 | Hs.88162 | Hs.86619 | Hs.85015 | Hs.84630 | Hs.83938 | Hs.82669 | Hs.81796 |
| Gene Name | | | | | | | | |
| Genbank | AW007080 | AA927475 | Al913396 | AA480075 | Al417267 | Al288745 | Al346341 | A1123555 |
| Affy | 47138_at | 62974_at | 48268_at | 62952_at | 55077_at | 91320_at | 62942_at | 54593_at |
| SeqID | 921 | 256 | 022 | 131 | 462 | 415 | 431 | 351 |
| # | 403 | 404 | 405 | 408 | 407 | 408 | 409 | 410 |

| | | | 9 | 1 | | | |
|---------------------------|---|-------------|---|--|---|--|-------------|
| p-values 1.8411E-07 | 5.4545E-06 | 2.739E-06 | 0.00000603 | 0.00000139 | 6.8391E-05 | 7.2051E-06 | 5.4948E-09 |
| Fold Change 0.29962365 | 0.280798539 | 3.262260583 | 0.27595812 | 0.277326235 | 3,506341539 | 0.29408728 | 0.222082398 |
| | Cluster Incl. Al692878:wd42c12.x1 Homo sapiens cDNA, 3 end flone=IMAGE-2330806 Alchone_and=3' flgb=Al692878 fgi=4970218 Alg=H8.78605 flen=556 | | Cluster Ind AL031846:aJ742C19.5 (novel Chromobox protein) /cds=(89,844) /gb=AL031846 /gj=4164388 /ug=Hs.7442 /len=364 | Cluster Incl AF070648:Homo sapiens clone 2485f mRNA sequence Icds=UNKNOWN igb=AF070648 (gi=3283922 /ug=Hs.74034 Alen=1313 | Cluster Incl. AW003215:wq64b02.x1 Homo saplens cDNA, 3 end /ctone=INAGE-2476011 /ctone_end=3 /gb=AW003215 /gl=8850131 /ugHs.73452 lan=655 | Cluster Incl. N95620:zb66b09.sr Homo saptens cDNA, 3 and /clone=IMAGE-308537 /clone_end=3' /pe=N95620 /gl=1267890 /ug=Hs,7212 /len=563' | |
| Cluster# Hs.7913 | Hs.76605 | Hs.76550 | Hs.7442 | Hs.74034 | Hs.73452 | Hs.7212 | Hs.72089 |
| Gene Name | | | | | | | |
| Genbank W73230 | Al692878 | AA046853 | AL031846 | AF070648 | AW003215 | N95620 | AA775711 |
| Affy 54992_at | 64747_at | 46274_at | 36894_at | 36119_at | 55986_at | 57214_at | 52844_at |
| SeqID 1203 | 601 | 27 | 853 | 304 | 910 | 1099 | 221 |
| # 1- | 412 | 413 | 414 | 415 | 416 | 417 | 418 |

| 110 02/00/ | | | 9 | 2 | | 101/ | |
|---|---|--|-------------|-------------|-------------|------------|---|
| p-values 0.00059258 | 0.00037428 | 5,5293E-06 | 2.0222E-07 | 2.0661E-06 | 3.7787E-05 | 1.4531E-07 | 0.00033754 |
| Fold Change 0.272334447 | 3.317504451 | 0.313288626 | 0.311833232 | 3.820099432 | 0.307655933 | 0.20785966 | 0.321089692 |
| Cluster Description Cluster Incl. AA142875:z499b06:s1 Homo sapiens cDNA, 3 end Idonne-IIMAGE-616535 hole, end-3" (pb=AA142875 (g=11112261 Iune-te, 71719 Nent-450 | Cluster Incl. AI742057.wg38d12.xi Homo sapiens cDNA, 3 end /done=IMAGE-2367383 /done_end=3 /gb=AI742057 /gi=5110345 /ureHs.7155 en=503 | Cluster incl. R51371;yg/7803.s1 Homo saplens 0.3132886226 CDNA; 3 end Clotne=IMAGE-38107 CAOLe_and=3'(pb=R51371 /gj=813273 Aug=1s,7107 Aerr=54.2 | | | | | Cluster Incl. Al948551:wp91c07.x1 Homo sapiens cDNA, 3 end /clone=IMAGE-2489132 /clone_end=3'/gb=Al948551 /gj=5740861 /ug=Hs.67317 /len=434 |
| Ctuster#*** Hs.71719 CI ss (ol | Hs.7155 | Hs.7107 | Hs.71058 | Hs.70704 | Hs.68505 | Hs.6750 | Hs.67317 |
| Gene Name | | | | | | | |
| <u>Genbank</u> AA142875 | Al742057 | R51371 | Al694389 | AA056180 | AI791751 | AL039870 | Al948551 |
| Affy 53762_at | 46659_at | 64913_at | 46649_at | 65999_at | 53733_at | 46622_at | 53724_at |
| SeqID 56 | 628 | 1110 | 905 | 58 | 999 | 861 | 799 |
| # 418 | 420 | . 451 | 422 | 423 | 424 | 425 | 426 |
| | | | | | | | |

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| WO 02/05927 | 1 | | 93 | | | PCT/ | JS02/02176 |
|--|---|---|---|------------|--|---|-------------|
| p-values 2.84E-04 | 6.5166E-08 | 0.00238804 | 1.7127E-07 | 1.3008E-05 | 0.00046176 | 0.00039039 | 7.743E-06 |
| <u>Fold Change</u> 0.327537441 | 0.215531153 | 3.179888739 | 0.262677342 1.7127E-07 | 0.09523078 | 0,311206678 | 0.309546056 | 0.305213649 |
| Cluster Ind ALGOSOF-Homo sophers mRNA; cDNA DKFZP664A026 (from done DKFZP664A026 (from done DKFZP664A026) (ASS=UNKNOWN AGP=ALGOSOF (pi=4914600 /ug=1s, 66762 | netr-20 Cluster Ind. AW028241.wv10d12.x1 Homo sapiens cDNA, 3 end /done=IMAGE-990854 folone_end=9.fgh=AW028241/gl=9878771 hiverlet 6x330 Ann=x30r | To this property of the prope | Cluster Incl. Al640524.wa29b02.x1 Homo sapiens cDNA, 3 end /clone=II/A/GE-229467 /clone=md=3 (gb=4640524 /gi=4703633 /nc=45.8382 /len=471 | | Cluster Incl. W68034;zd39e02.r1 Homo sapiens cDNA, E end /clone=IMAGE;43034 /clone_and=5 /gb=W68034 /gi=1376903 /kic=14s.6052 /len=533* | Cluster Incl. A4723692:ah85c11.s1 Homo sapiens CDNA, 3 end /clone=INA/GE-1325876 /clone ender3 (pla-A723682 /gi=2741399 /incl-Hs 5899 Incr-4911 | |
| Cluster # | Hs.65239 | Hs.6459 | Hs.6382 | Hs.62492 | Hs.6052 | Hs.5889 | Hs.58606 |
| Gene Name | | | | | | | |
| Genbank AL050367 | AW026241 | A1092936 | Al640524 | AA742697 | W68034 | AA723692 | A149693 |
| Affy 36821_at | 53687_at | 54001_at | 46583_at | 46200_at | 62493_at | 53200_at | 64720_at |
| SeqID 893 | 949 | 346 | 544 | 214 | 1193 | 212 | 373 |

| | | 110 02 000 | | | 9. | 4 | | | |
|---|-----------------------|--|--|---|---|---|---|--|--|
| | p-values | 0.00051417 | 4.6226E-06 | 6.2106E-06 | 7.5477E-08 | 1.5685E-09 | 2.5792E-05 | 1.2863E-05 | 4.7788E-05 |
| | Fold Change | 0.241273698 | 0.206509126 | 0.168279383 | 0.179788238 | 0.068952382 | 0.28516587 | 0.323048827 | 0.274783293 |
| | - Cluster Description | Cluster Incl. W73386:zd53e05.s1 Homo sapiens cDNA, 3 end /clone=INA/GE-344384 /clone_end=7 /gb=W73386 /gi=1383519 /ug=Hs. 58303 /fen=418' | Cluster Incl. AA044828:zk72c09.s1 Homo sapiens cDNA, 3 end (donne-IINAGE-488368 felone-ende-3' (gb-8A044828 /gi=1523031 /ug=Hs.86043 /fen=641' | Cluster Incl. AA166620:zo85f08.s1 Homo sapiens cDNA, 3 end /slone=INAGE-593703 /clone_end=3'/gb-44166520 /gi=1745209 /kg=Hs.55778 /len=697' | Cluster Incl. AA036952;zx30h01,s1 Homo saptiens CoDNA, 3 end /done=INA/GE472081 /done_end=3'/gb=AA036952 /gi=1510009 /ug=Hs.5084 / fan=571' | Cluster Incl. AI799784:wc43b08.x1 Homo sapiens CDNA, 3 end /clone=INA/GE-2321367 /clone_end=3' /gb=AI799784 /gi=5365256 /kg=Hs.49696 Ain=382' | Cluster Incl. AI743925:wg54f04.x1 Homo sapters CDINA, 3 end /clone=INA/GE-2368927 /clone_end=3'/gb=AI743925 /gi=5112213 /ug=Hs.4944 /len=505' | Cluster Incl. AA243659-zr68c06.s1 Homo sapiens CDNA, 3 end /clone=IMA/GE-688654 /clone_end=3' /gb=AA243659 /gi=1874478 /ug=Hs.4863 /len=481' | Cluster Incl. AA082546;ze88h10.r1 Homo sapiens cDNA, 5 end /clone=IMAGE-366115 /clone_end=5'/gb=AA082546 /gi=1624603 /ug=Hs.48516 /len=5677 |
| í | Cluster# | Hs.58303 | Hs.58043 | Hs.55778 | Hs.50841 | Hs.49696 | Hs.4944 | Hs.4863 | Hs.48516 |
| | Gene Name | | | | | | | | |
| | Genbank | W73386 | AA044828 | AA166620 | AA036952 | AI799784 | Al743925 | AA243659 | AA082546 |
| | Affy | 46087_at | 54407_at | 62213_at | 53011_at | 64694_at | 52986_at | 52183_at | 45786_at |
| | SedID | 1204 | 8 | 74 | 6 | 678 | 637 | 88 | 37 |
| | ## | 435 | 436 | 437 | 438 | 439 | 440 | 144 | 442 |
| | | | | | | | | | |

| WO 02/059 | 9271 | | 95 | | | PCT/U | US02/02176 |
|--|--|--|-------------|---|-------------|------------|--|
| p-values 2.6745E-08 | 0.00013182 | 5.9049E-07 | 1.5186E-06 | 4.1704E-06 | 7.1664E-06 | 6.2406E-11 | 6.5731E-05 |
| Fold Change 0.074704469 | 0.237704503 | 0.10436228 | 0.279947233 | 0,324029566 | 0.276986241 | 0.07793742 | 0.298567674 |
| Cluster Incl. Al200456:qf93003.xf Homo sapiens cDNA, 3 and dome=IMAGE-1757573 (done_end=? tgb=Al200456 (gj=3753062 hom=2A77) | og istactions and Constitution and Const | Cluster Inc. ALO4694:10FFZp58810717_r1 Homo sapiens cDNA, 5 end Clone=DKFZp58810717 (slone_end=5) (gb=AL046941 (gi=5435000 /ug=Hs.48531 An=704 | | Cluster Inci. N22378;yw37d04.s1 Homo sapiens cDNA, 3 end (clonne-IINA/GE-254407 Iclone_end=3' (gb=N22378 (gi=1128612 NucHs.43157 / Ann=450 | | | Cluster Incl. AI6512.12.wa98a10.x1 Homo sapiens cDNA, 3 end /clone=IMAGE-2304186 /clone_end=3'/gb=AI651212/gi=4735191 /ug=Hs.4283 /len=609' |
| Cluster# Hs.48516 Cl | Hs.47447 | Hs.46531 | Hs.44038 | Hs.43157 | Hs.43148 | Hs.43125 | Hs.4283 |
| Gene Name | | | | | | | |

AA150501

45220_at

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448

N22378

1065 59014_at

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AI864898

52080_at

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449

AI651212

52075_at

548

450

Genbank Al200456

Affy 51012_at

SeqID 383

44

AA541622

64407_at

162

444

AL046941

52140_at

876

445

AI768516

62136_at

658

| | | | | | : | 96 | | | |
|---|---------------------|--|---|------------------------|--|-------------|---|-------------|---|
| | p-values | 0.00148036 | 0.00152421 | 1.1279E-08 | 1.5134E-05 | 3.4237E-06 | 1.848E-05 | 6.9772E-05 | 3.554E-07 |
| | Fold Change | 0.317296872 | 3.149001267 | 8.270850261 1.1279E-08 | 3.94 | 0.276493253 | 5.223054257 | 3.048509737 | 4.526523002 |
| | Cluster Description | Cluster Incl. A1934361:wp04g11.xt Homo saplers 001Nq, 3 and /clone=INIAGE-2463908 /clone_end=3'/gb=A184361/gl=5673231 /ug=Hs.42886 flep=688* | Cluster Ind. Al990483:ws40b12.x1 Homo saplens cDNA, 3 end /done=II/NAGE-2499647 /done_end=3 /gb=Al990483 /gi=5837364 /ug=Hs.4243 /hen=541 | | Cluster Incl. AI751438:cn10a03.y1 Homo sapiens cDNA (clone=NHTBC, cn10a03-(random) (gb=AI751438 (gb=5129702 /ug=Hs, 162827 Hen=513 | | Cluster Incl. N25267;yy74h01.s1 Homo saplens cDNA, 3 end /clone=INA/GE-267505 /clone_end=3 /gb=N25267 /gl=1139417 /uz=Hs.40479 /len=460' | | Cluster Ind. A4921830:om44b04.st Homo sapiens cDNA, 3 end /clone=IMAGE-1543855 /clone_end=3'' /gb=A4921830 /gl=3069139 /ug=Hs.38178 /len=516' |
| , | Cluster# | Hs.42586 | Hs.4243 | Hs.41271 | Hs.41271 | Hs.41067 | Hs.40479 | Hs.39421 | Hs.38178 |
| | Gene Name | | | | | | | | |
| | Genbank | Al934361 | A1890483 | AI557210 | AI751438 | AL079707 | N25267 | W89022 | AA921830 |
| | Affy | 45779_at | 45203_at | 52019_at | 85126_at | 45179_at | 46372_at | 46365_at | 51970_at |
| | SedID | 787 | 846 | 512 | 149 | 968 | 1068 | 1212 | 254 |
| | #1 | 451 | 452 | 453 | 454 | 455 | 456 | 457 | 458 |

| WA | 02/ | 050271 |
|----|-----|--------|

| WO 02/059271 | | | 97 | | PCT/US02/02176 | | |
|--|--|--|---|------------|----------------|--|--|
| p-values 2.3349E-06 5.5911E-06 | 2.893E-08 | | 0.00164325 | 1.2131E-05 | 3.3983E-05 | 0.00080689 | |
| 6.25714791 0.25714791 0.279922681 | 4,913802444 | 3.065154029 0.01073513 | 0.306446993 | 0.33 | 3.165598561 | 0.310421016 | |
| Cluster Incl. A70997 Suvedenza xt Homo Superior DNA, a and Johne-IMAGE-2221893 Include and-3 (git-A17997 In git-9235448 Ingerles 38 Fto Name-547 Cluster Incl. A806221 wt/2809 xt Homo Superior SDNA, 3 and Johne-IMAGE-2385744 Incline, gend-2 (git-8808271 /gjt-8892787 Incline, gend-2 (git-8808271 /gjt-8892787 | mg-11s.00cz nat-0c Cluster Incl. W02608zas51g08.r1 Homo sapiens cDNA, 5 end /clone=IMAGE-296126 fictore_end=5 (gb=V02608 /gj=1274588 hin=He 36301 /hn=HR | The state incl. H71532:ys11909.s1 Homo sapiens cDNA, 3 end /clone=IMAGE-214528 /clone_end=3 /gp=17332 /gj=1043348 /nr=H5 36923 /mn=222 | Cluster Incl. A4/79633:m32d12.x1 Homo sapiers cDNA, 3 end /clone=IMAGE-2158295 (Ichone-end-2) (pab-4479633 (gl=4372801 incl+3.3716 /nen=489 | | | Cluster Incl. W73855zd5Zf10.r1 Homo sapiens 0.310421016 cDNA, 5 end (done=IMAGE-544299 lobone_end=5' (gb=W73855 (gi=1384028 lug=Hs.32343 Aere577 | |
| Cluster# Hs.38163 Hs.38022 | Hs.36830 | Hs.36823 | Hs.33716 | Hs.32615 | Hs.32458 | Hs.32343 | |
| Gene Name | | | | | | | |
| Genbank Al799976 Al806221 | W02608 | H71532 | Al479633 | AA830307 | AI808983 | W73855 . | |
| Affy 65094_at 49825_at | 61727_at | 60143_r_at | 64252_at | 88308_at | 45896_at | 45757_at | |
| 679 679 686 | 1170 | 982 | 485 | 234 | 689 | 1206 | |
| 459 460 460 | 461 | 462 | 463 | 464 | 465 | 466 | |

| | WO 02/059 | WO 02/059271 98 | | | | | | JS02/02176 |
|--------------------|---|--|-------------|---|-------------|--|---------------------|---|
| aouley-d | 4.1587E-06 | 5.9115E-06 | 1,4632E-05 | 0.00063822 | 0.00321371 | 0.00228259 | 0.00321198 | 0.00016645 |
| Fold Change | 0.286699064 | 0.171580051 | 0.196081354 | 0.287881302 0.00063822 | 0.324205584 | 0.227579801 | 0.309546276 | 0.283008853 |
| Olitetor Departmen | Cluster Incl. / sapiens cDN /clone_end=3 /ug=Hs.3224 | Cluster Incl. W07043:za92e08.r1 Homo sapiens cDNA, 5 and /clone=INA/GE-300038 /clone_end=6 /gb=W07043 /gi=1281065 /uc=Hs.23156 /len=692" | | Cluster Incl. AA921922:com40h06.s1 Homo sepiens cDNA, 3 end /clone=INA/GE-1643647 /clone_end=3 /gb=AA921922 /gi=3069231 /ug=Hs. 31412 /len=489* | | Cluster Incl. AI743715:wg63a08.x1 Homo sapiens cDNA, 3 end /clone=In/AGE-2368790 /clone_end=3 /gb=AI743715 /gj=5112003 /uzeHs.37359 /len=489 | | Cluster Incl. Al093702:qe31f05.s1 Homo sapiens cDNA, 3 end /clone=IMAGE-1688385 /clone_end=3'/gb=Al093702 /gj=3432678 /ug=Hs. 169169 /len=358' |
| O. rotor # | Hs.32241 | Hs.32135 | Hs.31695 | Hs.31412 | Hs.31297 | Hs.31297 | Hs.30835 | Hs.30156 |
| Comp Money | | | | | | | | |
| Jacque | Al913749 | W07043 | AL040912 | AA921922 | AI557360 | Al743715 | AA195251 | Al093702 |
| | 44679_at | 64238_at | 44575_at | 61681_at | 43591_r_at | 50955_at | 65651_s_at AA195251 | 50658_s_at |
| 2 | 177 | 1173 | 365 | 255 | 513 | 636 | 88 | 347 |
| 4 | 467 | 468 | 469 | 470 | 471 | 472 | 473 | 474 |

| | | | | 9 | 9 | | | |
|---------------------|---|-------------|-------------|------------|-------------|-------------|-------------|---|
| p-values | 5.309E-12 | 7.704E-05 | 0.0041327 | 0.00030044 | 2.469E-12 | 0.00021707 | 1.3716E-06 | 4.6046E-08 |
| Fold Change | 13.96572736 | 0.274066497 | 0.321891202 | 0.20 | 4.243716901 | 0.259103459 | 0.211477764 | 0.131870144 |
| Cluster Description | Cluster Incl. AA059458:z198g05.r1 Homo sapiens cDNA, 5 end /clone=INA/GE-512504 /clone_end=5 /cp=AA059458 /gi=1553317 /un=Hs_28792 /len=572 | | | | | | | 6. Cluster Incl. A6805411w82c09.x1 Homo sapiens cDNA, 3 end /done=IMAGE-2286192 /done_end=3' /gb=A680541 /gj=4890723 /ug=Hs_25173 /len=1072* |
| Cluster# | Hs.28792 | Hs.28625 | Hs.286228 | Hs.286216 | Hs.286117 | Hs.285995 | Hs.285966 | Hs.285966 |
| Gene Name | | | | | | | | |
| Genbank | AA059458 | Al806324 | Al439628 | H54254 | AA705188 | AA126704 | AA131648 | Al680541 |
| Affy | 65626_at | 44025_at | 91773_at | 78617_at | 53490_at | 52999_at | 56211_at | 50408_at |
| SeqID | 8 | 687 | 474 | 992 | 206 | 14 | 64 | 588 |
| #1 | 475 | 476 | 477 | 478 | 479 | 480 | 481 | 482 |

| WO 02/03/2/1 | | | | . 1 | 00 | | 101/0002022 | |
|-------------------------|--|--|-------------|--|---|------------|---|--|
| p-values | 0.00017459 | 0.00011017 | 0.00024096 | 0.0008652 | 6.1606E-11 | 0.00027317 | 5.8287E-05 | 5.0013E-05 |
| Fold Change | 0.30 | 0.262417057 0.00011017 | 3.603536057 | 0.327647494 | 11.21387388 | 0.18 | 0.326231649 | 0.22 |
| Cluster Description | Hs.286834 Cluster Incl. Al656062:tt43b04.x1 Homo sapions cDNJ, and klome=IMAGE-2243603 klone_end=3' kgb=Al656062 kj=4740041 kig=Hs.293724 len=526 | Cluster Incl. W02823:za65h04.r1 Homo sapiens cDNA, 5 and /clone=IMAGE:291703 /clone_end=5 /gb=W02823 /gi=1274868 /ug=16.234968 /len=461* | | Cluster Ind. W19285;zb90g08.s1 Homo sapiens cDNA, 3 end /done=IMAGE:310910 /done_end=3' /gb=W19285 /gi=1294973 /ug=Hs,202731 /len=460' | Cluster Incl. AA669106:aa81g05.s1 Homo saplens cDN4, 3 end /donn=IMA/GE-827384 /done_end=3'/gb=AA699106 /gi=2630605 /ug=Hs,108106 /len=637* | | Cluster Incl. A1963873:wt86b08.x1 Homo sapiers cDNh, 3 and /olone=INAGE-2614327 /clone_and=3 /gb-A1963873 /gi=5756586 /ug=Hs.227082.len=704 | Cluster Incl. Al557450:PT2.1_T_D12.r Homo sapiens cDNA, 3 end /clone_end=3' /gb=Al557450 /gj=4489813 /ug=Hs.235996 /len=565' |
| Cluster# | Hs.285834 | Hs.285785 | Hs.285590 | Hs.285570 | Hs.285473 | Hs.285414 | Hs.285247 | Hs.285233 |
| Gene Name | | | | | | | | |
| Genbank | Al656062 | W02823 | Al458306 | W19285 | AA669106 | Al983045 | Al963873 | Al557450 |
| Affy | 88239 at | 50990_at | 59070_at | 65988_at | 48083_at | 87998_at | 60842_at | 78103_at |
| SedID | 228 | 1171 | 478 | 1175 | 198 | 838 | 815 | 514 |
| #1 | 483 | 484 | 485 | 486 | 487 | 488 | 489 | 490 |
| | | | | | | | | |

| | | | | | , 1º | 01 | | | ., |
|---|---------------------|--|---|---|-------------|---|------------|---|--|
| | p-values | 1.8665E-06 | 0.00150517 | 8.0203E-05 | 3.4232E-05 | 8.7733E-05 | 0.00121855 | 2.5913E-05 | 0.00082565 |
| | Fold Change | 4.466366979 | 0.31 | 0.254762986 | 4.911742129 | 0.27 | 0.23 | 0.30 | 4.06 |
| | Cluster Description | Hs.285220 Cluster Incl. A869951⋅wi63a07.x1 Homo sapplems cDNA, 3 and idone=INAGE-2A29556 idone_and=3' (gb=4869951 (gj=5543919 iug=Hs.2085 fleer=752. | 9 Cluster Ind. N31046;yx51h06.r1 Homo sapiens cDNA, 5 end /done=IMAGE-265307 /clone_end=5 /gb=N31046 /gi=1151445 /ug+ls.154508 /en=422* | Otuster Incl. AA824349:aj30f10.s1 Homo sepiens cDNA, 3 end /clone=1391851 /clone_end=3' /gb=AA824349 /gi=2896419 /ug+Hs 40300 /len=888* | | O Cluster Incl. A4435443:th94e06.x1 Homo sapiens cDNA, 3 end (done=INAGE-2126338 /clone_end=3' /gb=A435443 /gi=4303218 /urg+ls. 149064 len=475 | | 4 Cluster Incl. A188749:qd11c05.x1 Homo sapiens cDNA, 3 end (clone=IMAGE-1723400 Rdone_end=3' (pl=A188749 (gl=3739958 lug=ls, 181742, Ien=467 | O Cluster Incl. Al446168:ij07h07.x1 Homo sapiens cDNA, 3 end /clone=IMAGE-2140861 /clone_end-a"/gb=Al446168 /gj=4283320 /ug=Hs.176708 /len=435 |
| ļ | Cluster# | Hs.285220 | Hs.285009 | Hs.284269 | Hs.283713 | Hs.276860 | Hs.274252 | Hs.271594 | Hs.271530 |
| | Gene Name | | | | | | | | |
| | Genbank | Al869951 | N31046 | AA824349 | AA584310 | Al435443 | AA143491 | Al188749 | Al446168 |
| | Affy | 63460_at | 83506_at | 51999_at | 48774_at | 83118_at | 78658_at | 67167_at | 84893_at |
| | SeqD | 749 | 1070 | 232 | 171 | 470 | 8 | 379 | 476 |
| | ## | 191 | 492 | 493 | 494 | 495 | 496 | 497 | 498 |

| | | WO 02/059 | 271 | | 10 | 12 | | PC | T/US02/02176 |
|----|-------------------------------|---|---|--|---|--|---|--|---|
| | b-values | 0.00710567 | 0.01228173 | 5.7601E-05 | 9.3673E-05 | 0.00017276 | 0.00654385 | 0.01366367 | 0.00854584 |
| | Fold Change | 0.315940639 | 3.07 | 0.18 | 4.593843245 | 0.250833383 | 0.247069023 | 4.683322065 | 0.28 |
| i- | Cluster # Cluster Description | Hs.271363 Cluster Incl. Rt17937;ye90706 at Homo sapiens 0.315940639 CBNA, 3 end follone=MAGE-175927 Id=771547 Inclone_end=3 (pa=Rt17937 ig=771547 Iug=Hs.67734 ip=Rt248 Inclone_end=3 inclone_end=3 inclone=3 | Hs.277157 Cluster Incl. AA677864:zi13d12.s1 Homo sepiens cDNJ, 3 end (donne=INAAE=430679 /clone_end=3'/gp=AA677864 /gi=2658386 /ug=Hs.222705 flen=384 | Hs.270549 Cluster Incl. W72407.zd67c03.sr1 Homo sapiens cDNA, 3 end cloone=INAGE-345700 /clone_end=3'/gb=W72407 /gl=1382424 /ug=Hs: 118607 /len=651' | Hs.270524 Cluster Incl. A1743516:wif72b10.x2 Homo sppiens cDNA, 3 end clonne=INAGE-2361115 /clone_end=3'/gp=A1743516 /gi=5111804 /ug=Hs.205320 /len=683 | Hs.270235 Cluster Incl. Al418596:tg37d02.x1 Homo sapiens cDNA, 3 and iclone=INIAGE-2110947 iclone_end=3'fgb=Al418596 kgi=4264527 kgi=Hs; 187926 ken=43.1 | Hs.270027 Cluster Incl. AI683911:tw54f08.x1 Homo sapiens cDNA, 3 end /donne=INIA/GE-2263827 /done_end=3'/gb=AI83911 /gi=4894093 /ug=Hs: 445791 /len=497 | Hs.269628 Cluster Incl. AA773348:ab56904.s1 Homo sapiens cDN0, a rend Idonne=845718 /clone_end=3'/gb=AA773348 /gi=2624919 /ug=Hs.192544 /len=608 | Hs.269392. Cluster Incl. R69564/yi40x04.st Homo sapiens OLMA, 3 end Indone=IMAGE-141726 Idone, part—2' (pl=-R6956 kjg=843101 /ug=+16.183359 / Ren=379' |
| | Gene Name | _ | _ | - | _ | _ | _ | _ | _ |
| | Genbank | R17937 | AA677864 | W72407 | AI743516 | Al418596 | Al683911 | AA773348 | R69584 |
| | Affy | 60202 <u>i a</u> t | 72092 <u>f</u> at | 76208_at | 69687_at | 87016_at | 91206_at | 88243_r_at | 78883_at |
| | SedID | 1103 | 199 | 1200 | 633 | 466 | 591 | 219 | 1115 |
| | ## | 499 | 200 | 501 | 502 | 503 | 504 | 505 | 506 |
| | | | | | | | | | |

| | P | CT/U | S02/0 | 2176 | |
|---|---|---|--|---|--|
| | 0.00127676 | | 0.01815078 | | |
| | 3.03 | | 0.310940166 | | |
| /cione_end=3' /gb=AI/42002 /gl=5110210 /ug=Hs.239805 /len=490' | Hs.250594 Cluster Incl. Al672389:ty64f02.x1 Homo sapiens cDNA, 3 end /clone=IMAGE-2283867 | /cione_end=> /gb=Anb/2503 /gr=4632120 /ug=Hs.139891 /len=587 | Hs.24898 Cluster Incl. R67627:yj69c11.s1 Homo sapiens 0.310940166 0.01815078 cDNA, 3 end /clone=IMAGE-154004 | /done_end=3'/gb=R67627 /gi=840265 /ug=Hs.24898 /len=474' | |
| | Hs.250594 | | Hs.24898 | | |

| | WO 02/059 | 271 | | 103 | | | PCT/U | JS02/02176 |
|-------------------------------|--|--|---|--|--|---|--|--|
| p-values | 9.1918E-06 | 3.1637E-07 | 7.5535E-05 | 0.00045064 | 6.5167E-06 | 0.00715494 | 0.00127676 | 0.01815078 |
| Fold Change | 0.142950221 | 0.27 | 0.092486133 | 0.194270285 | 0.27933205 | 3.052533662 | 3.03 | 0.310940166 |
| Cluster # Cluster Description | Hs.268286 Cluster Incl. R20784;)h18b08.s1 Homo sapiens cJNN, a send icono=INIAGE-130071 icono=INIAGE-130071 icono=INIAGE-130071 icono=INIAGE-130071 icono=INIAGE-130071 icono=INIAGE-130071 icono=INIAGE-130071 icono=INIAGE | Hs.265499 Cluster Incl. AW014647/UI-HBI0p-abd-b-T2-O- ULS1 Home sapiens cDN4.2 and convenience of idenoral MAGE-Z711475 (done_and-a') idenoral MAGE-Z711475 (done_and-a') iden=402' iden=402' iden=402' iden=402' iden=402' | Hs.260164 Cluster Incl. R535943/j71c01.s1 Homo sapiens cDNA, 3 end clonne=IIIAAGE-154176 Idone_and=3' /gp=R53594 /gj=815496 ing=Hs.27444 Ilen=468 | Hs.25391 Cluster Incl. Af524085;th01e09.x1 Homo sapplens cDNA, 3 end (clone=INAGE-2117032 /clone=Ind=3')gb=Af524085 /gi=4438220 /cjg=Hs.2539 /len=509" | Hs.25248 Cluster Incl. A1659533:tu12a12.x1 Homo saplens GDNA, and Idone=INIAGE-2260814 Ichone_end=3' (pb=41659633 /gi=4763103 Iup=Hs.25248 Ilen=654' | Hs.250879 Cluster Incl. A1742002.wg37f06.x1 Homo sapiens cDNA, 3 end idono=INIAGE-2367299 / klone_end=9 / gp=A1742002 /gi=51.10210 / klcr=1s.239805 /en=490 | Hs.250694. Cluster Incl. AI672389:ty64f02.x1 Homo sapiens cJNA, 3 end clonne=II/IA/GE_2283867 / clone_end=3'/gb=Ai672389 /gi=4852120 / cg=Hs.1398091 /len=587. | Hs.24996 Cluster Incl. R67627/3/69c11.s1 Homo saplens 0.310940166 CDNA, 3 end /clone=IMAGE-154004 Clone ender-3/pmeR6727 /gj=640265 Clone ende |
| Gene Name | _ | _ | - | | | - | - | |
| Genbank | R20784 | AW014647 | R53594 | Al524085 | Al659533 | AI742002 | Al672389 | R67627 |
| Affy | 49549_at | 80401_at | 42913_f_at | 64057_at | 50411_at | 61333_at | 78487_at | 51886_at |
| SedID | 1104 | 932 | 111 | 200 | 566 | 627 | 578 | 1114 |
| ## | 507 | 208 | 909 | 210 | 111 | 512 | 513 | 414 |

| wo | 02/059271 | | | 104 | | PC | CT/US02/02176 |
|--|---|-------------|--|-------------|---|--|---------------|
| p-values 0.00047742 | 1.293E-05 | 2.1691E-05 | 7.4615E-08 | 1.1547E-07 | 3.2473E-05 | 4.0505E-06 | 0.00326112 |
| Fold Change 0.247364362 | 0.280733599 | 0.324185103 | 3.482682058 7.4615E-08 | 0.227760861 | 0.264360188 | 0.221960648 | 0.30 |
| Cluster Description Cluster Incl. AA045145.2711412.r1 Homo saniens cDNA 5 end (chon=INAGE-37683) | Adone_end=5' (pb=AA045145 /gi=1623485 Ag=Hs.24872 Aen=588' Cluster Incl. AT763199wg35b03.x1 Homo senians cPINA.3 and Adone=IMAGE-2487053 | | (clone_end=5'(pb=A4/27/27')gi=1687016 hg=18_24715 flen=611' Cluster Inci. AA224205:zr15f03.s1 Homo sepiers cDINA, 3 end /clone=IMAGE-683486 inchen end=5'(hp=A42/205f cln=184747 | | Nug=Hs.19251b Inter=440 ¹ Cluster Incl. Al658662:Lu22a11.x1 Homo sapiens cDNA, 3 end Idone=IMAGE-2261772 Idone_end=3' (µb>Al658662 /qi=4762232 | Cluster Inc. AA034289:zk18e07.r1 Homo sapiens cDNA, 5 end /clone=IMAGE-470916 idnose ander5f (gp=4034289 /gj=1506098 hursts ondard AnderseR | |
| Cluster# Hs.24872 | Hs.24790 | Hs.24715 | Hs.24529 | Hs.243010 | Hs.24192 | Hs.24192 | Hs.241797 |
| Gene Name | | | | | | | |
| Genbank AA045145 | AI769199 | AA127727 | AA224205 | AI583530 | Al658662 | AA034289 | AA909818 |
| Affy 43919_at | 43554_at | 50385_at | 50018_at | 65523_at | 50361_at | 60532_at | 77970_at |
| SeqID 24 | 099 | 84 | 88 | 522 | 561 | 17 | 248 |
| # 212 | 516 | 517 | 518 | 519 | 520 | 521 | 522 |

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| WO 02/059271 | | | 105 | | | PCT/US02/02176 | | |
|--|--|---|-------------|------------------------|---------------------|------------------------|------------|---------------------|
| p-values 4.329E-05 | 1.5567E-06 | 2.5531E-05 | 9.8105E-06 | 4.0549E-05 | 1.9829 E- 05 | 0.00035218 | 0.00788877 | 1. 7239E-0 6 |
| Fold Change 0.313634138 | 0.232069434 | 0.31162811 | 4.191125642 | 0.325095547 4.0549E-05 | 0.24 | 3.938205017 0.00035218 | 0.20 | 0.290657123 |
| Cluster Incl. C16443:C16443 Homo sapiens cluNA, 5 end clone=CEN-221F12 clone_end=5' (pb=C16443 /gi=1571150 /un=Hs.2414 /len=456) | Cluster Ind. AI797276:we86f09:x1 Homo sapiens cDNA, 3 end /done=IMAGE-2348009 /done_end=3 /gb=AI797276 /gi=5362748 /irr=Hs 23912 Inn==516* | Cluster Incl. At/21837:tf56c10.x1 Homo sapiens cDNA, 3 and /clone=IMAGE-2103186 flotine_and=3 (pa-4/21837 /gl=4287768 | | | | | | |
| Cluster# Hs.24144 | Hs.23912 | Hs.23869 | Hs.237809 | Hs.23630 | Hs.235920 | Hs.235758 | Hs.235390 | Hs.234898 |
| Gene Name | | | | | | | | |
| Genbank C16443 | AI797276 | Al421837 | AA948319 | N57539 | AA629715 | Al362288 | W26589 | Al970898 |
| Affy 50001_at | 43502_at | 56624_at | 52615_at | 56574_at | 73233_at | 66131_at | 78622_r_at | 43427_at |
| SeqID 956 | 674 | 468 | 262 | 1086 | 190 | 438 | 1178 | 822 |

| | WO 02/059 | 271 | | 1 | 06 | | PC | T/US02/02176 |
|------------------------------|---|--|---|--|---|--|--|--|
| p-values | 1.2854E-06 | 8.7469E-08 | 1.2339E-07 | 6.6059E-07 | 4.4094E-06 | 0.00181461 | 0.01016812 | 2.4083E-06 |
| Fold Change | 0.268475458 | 4.250714876 | 3,197888571 | 3.85742898 | 3.322756779 | 3.320734927 | 0.233986843 | 0.280053615 |
| Cluster# Cluster Description | 19,234898 Cluster Incl. A1057637.oy31h06.x1 Homo sapplens GDNA, 3 end Claone=INA/GE-1697483 /clone_end=9' (gb=A1057637 /gj=3331603 /ug=Hs,21305 flen=589' | Hs.234545 Cluster Incl. AI015982.ou95e08.x1 Homo sapiens cDNA, 3 and Idonp=INAGE-1635586 iclone_end=3' (gb=AI015982 gj=2230318 utu=1s, 59927) [en=485? | Hs.234545 Cluster Incl. AA383718:EST97358 Homo sapiens cDNA, 3 end clonne—ATOC-188064 clone_end=3'(gb=AA383718 /gj=2036227 Ng=Hs.234545 [len=311" | Hs.23448 Cluster Incl. A4133979.zzn88b09.r1 Homo sapiens aDNA, 5 end (done=IIMAGE-665241 /done_end=5' (gb=A4133979 /gj=1691065 /uc=Hs.22448 /len=569* | Hs.233634 Cluster Incl. Al970896;wr/21b11.x1 Homo sapiens cDNA, 3 end idono=IIIAAGE-2488317 idone_end=3'(gb=Al970896 igi=5767722 iug=1s.23364 Ilen=514 | Hs.233310 Cluster Incl. Al978650:wr67g08.x1 Homo saplens cINN., 3 end clonne=INAGE-2491838 /done_end="9" (pp=4978650 /gi=5803860 /ug=Hs.23310 /len=516 | Hs.232/17 Cluster Ind. Al208691:qg56b04.x1 Homo sapiens cbNA, 3 end idone=INAGE-1839151 (done_end=9/1gb=Al208691 (gl=3770633 idon=1942-23177 len=369 | Ha.23202 Cluster Incl. A4419030.tf63801.xr Homo aspiens cDNx, 3 and lobne=IMAGE-2102953 Iclone and-2º1 gip-A4419030 (pj=4284961 /ug=Hs.23202.flen=473 |
| Gene Name | _ | ; | _ | | _ | _ | _ | |
| Genbank | AI057637 | AI015982 | AA383718 | AA133979 | Al970896 | AI978650 | Al208691 | Al419030 |
| Affy | 49452_at | 55504_at | 64282_at | 44055_at | 44974_at | 74340_at | 74162 <u>r_</u> at | 43046_at |
| SeqID | 332 | 312 | 100 | 25 | 821 | 834 | 390 | 467 |
| ## | 532 | 533 | 534 | 535 | 636 | 537 | 238 | 539 |

| | | | | | 10 | 07 | | | |
|----|-----------------------|---|-------------------|------------|------------|------------|---|------------|-------------|
| | p-values | 7.5905E-05 | 8.6161E-05 | 1.3565E-05 | 5.2137E-05 | 0.00152718 | 9.4414E-05 | 0.00160759 | 0.00069879 |
| | Fold Change | 0.257584715 | 0.174616059 | 4.00 | 0.32 | 9.39 | 0.308687757 | 3.18 | 3.480820479 |
| | # Cluster Description | Cluster Incl. W45681.zxz86c02.r1 Homo sapters cDNA, 5 end /done=IMAGE-323426 /clone_end=5'/gb=W45681 /gl=1329681 /ug=He.23133 /fan=687* | | | | | Cluster Incl. AI816835:wj34712.x1 Homo saplens cDNA, 3 end /done=IMAGE:2404751 /clone_end=3' /gb=AI816836 /gi=5435914 /ug+Hs_222326 /len=526" | | |
| ٠. | Cluster # | Hs.23133 | Hs.22971 | Hs.227513 | Hs.225779 | Hs.224902 | Hs.222326 | Hs.222088 | Hs.220756 |
| | Gene Name | | • | | | | | | |
| | Genbank | W45581 | N91161 | AI829520 | Al956095 | Al499240 | AIB16835 | AA451665 | Al339240 |
| | Affy | 57119_s_at | 59769_s_at N91161 | 87583_at | 72501_at | 72363_f_at | 72674_at | 87339_at | 71899_at |
| | SedD | 1187 | 1097 | 730 | 808 | 497 | 701 | 123 | 425 |
| | ## | 540 | 143 | 542 | 643 | 544 | 545 | 546 | 547 |
| | | | | | | | | | |

| | WO 02/059 | 271 | | 10 | 08 | | PC | T/US02/02176 |
|-----------------------|---|-------------|-------------|-------------|------------|-------------|------------|--------------|
| p-values | 0.012637 | 7.7241E-05 | 0.00071317 | 1.0609E-07 | 2,5827E-06 | 3.9564E-05 | 0.00147634 | 0.00020743 |
| Fold Change | 3.51886622 | 3.531852021 | 0.306661245 | 0.085100991 | 0.2674744 | 0.286772796 | 0.22 | 0.30 |
| # Cluster Description | 19.220697 Cluster Incl. AA251131zs03b12.s1 Homo sapiens cDNA, 3 end clone=IMAGE-684095 /clone_end=v1 /gp=AA251131 /gj=1886093 /ug=Hs.2200897 /lenn=3697 | | | | | | | |
| Cluster# | Hs.22060 | Hs.22011 | Hs.21914 | Hs.218707 | Hs.218037 | Hs.214906 | Hs.21415 | Hs.213923 |
| Gene Name | | | | | | | | |
| Genbank | AA251131 | Al344312 | A1949833 | N63913 | Al911149 | AI610837 | A1935522 | AI917447 |
| Affy | 71839_at | 42988_at | 60813_at | 44210_at | 71668_at | 71524_at | 91345_at | 87161_s_at |
| SedID | 06 | 430 | 802 | 1090 | 765 | 532 | 791 | 477 |
| ## | 548 | 549 | 220 | 551 | 552 | 553 | 554 | . 555 |

| 110 02,005 | | | 1 | | 1,0002021,0 | | |
|---|---|------------------------|---|-------------|-------------|-------------|-------------|
| p-values 0.00795029 | 7.8969E-05 | 0.03460633 | 0.00010506 | 0.00648177 | 1.3875E-05 | 1.5821E-06 | 0.00062149 |
| Fold Change 0.292836533 | 3.74 | 0.302753776 0.03460633 | 3.62 | 3.047280178 | 0.23 | 6.957639593 | 3.289367551 |
| Cluster# Cluster Incl. AB88483wm32e0Z.xrl forno sapiens 60Mx, 3 and forne=INAGE-2447162 Idone_end=3 /gb=4889439 (gl=5589857 lughts.212709 len4.73 | Cluster Incl. AI823649:wi85g03.x1 Homo sapiens CDNA, 3 end /donne=IMAGE-2400148 /clone_end=3'/gb=AI823649 /gi=5444320 /tuchls_211535 flan=611' | | Cluster Ind. Al989871:ws38e12.x1 Homo sapiens cDN4, 3 end /donne=IMAGE-2499310 fclone_end=3' (gb=Al999871 /gj=5836752 /kg=15,210467 /len=722* | | | | |
| Cluster# Hs.212709 | Hs.211535 | Hs.211129 | Hs.210467 | Hs.209235 | Hs.209078 | Hs.208912 | Hs.208854 |
| Gene Name | | | | | | | |
| Genbank Al888493 | AI823649 | Ai825341 | Al989871 | Al800529 | Al351653 | Al381686 | H68822 |
| <u>Affy</u> 71153 <u>i</u> at | 87102_at | 70733_r_at Al825341 | 80045_at | 70350_at | 70704_i_at | 64145_at | 70219_at |
| SeqID 759 | 714 | 717 | 842 | 980 | 433 | 454 | 994 |
| 556 | 557 | 558 | 559 | 560 | 561 | 562 | 563 |

| | | WO 02/059 | 271 | | 110 | | | PC1/C | 1802/02176 |
|---|---------------------|---|--|-----------|---|------------|-------------|------------|------------|
| | p-values | 0.00825561 | 0.00066989 | 3.9E-05 | 5.8656E-06 | 0.00188255 | 7.7231E-05 | 4.6902E-05 | 7.3252E-09 |
| | Fold Change | 0.20 | 3.01 | 0.29 | 0.188450369 | 0.31 | 3.973692969 | 0.28 | 12.37 |
| - | Cluster Description | Cluster Incl. AI917901:tz14d04.x1 Homo saplers cDNA, 3 end /clone=IMAGE-2288551 /clone_end=3 /gb=AI917901 /gi=5637756 /ug=Hs.208641 /len=519* | Cluster Incl. AA628024:nq61f04.s1 Homo saplens cDNA, 3 end klonn=IMAGE-1148383 klone_end=3' kgb-aA628024 /gi=2540023 kug+1s.208588 klen=483* | | Cluster Incl. Al659783:tt86d01.x1 Homo sapiers cDNA, 3 end /clone=INA/GE-2288417 /clone_end=3 /gb=Al659783 /gl=4763353 /uc=Hs.207578 /len=452 | | | | |
| | Cluster# | Hs.208641 | Hs.208558 | Hs.208207 | Hs.207578 | Hs.205696 | Hs.205690 | Hs.204044 | Hs.203879 |
| | Gene Name | | | | | | | | |
| | Genbank | Al917901 | AA628024 | AL046628 | AI659783 | AI760534 | AI743599 | AI888991 | AI492879 |
| | Affy | 70637_at | 70631_at | 70592_at | 69893_at | 86820_at | 89203_at | 86750_at | 77842_at |
| | SeqID | 775 | 187 | 875 | 299 | 647 | 634 | 760 | 493 |
| | ## | 564 | 565 | . 286 | 292 | 268 | 269 | 929 | 571 |
| | | | | | | | | | |

| wo | 02/059271 |
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| | WO 02/059271 PCT/US02/02176 | | | | | | | | |
|---------------------|--|--|--|------------|------------|------------------------|--|--------------|--|
| | WO 02/059 | 2/1 | | 1 | 11 | | PC | 1/US02/021/0 | |
| p-values | 7.7642E-05 | 0.00187143 | 0.00030428 | 0.00090022 | 0.01216522 | 2.9782E-06 | 0.00527403 | 0.00084162 | |
| Fold Change | 4.96 | 3.17 | 3.64 | 0.29 | 4.11 | 0.263799925 2.9782E-06 | 0.300766236 | 0.322154385 | |
| Cluster Description | Cluster Incl. Al700646:we38h07.x1 Homo saplens cDNA, 3 end /clone=INAGE-2343421 /clone_end=3'/gb=Al700646 /gi=4986546 /ug=Hs.202269 /len=466 | Cluster Incl. Al937060:wp72f01.x1 Homo sapiens cDNA, 3 end /clone=INAGE-2487321 /clone_end=3'/gb=Al937060 /gi=5675930 /ug=Hs.202040 /len=522 | Cluster Incl. AI825713:wb75g02.x1 Homo septens cDNA, 3 and /clone=INAGE-2311538 /clone_end=3 /gb=AI825713 /gi=5446384 /ug=Hs.201975 /len=445 | | | | i Cluster Incl. Al653487.tq94h03.x1 Homo sapiens cDNA, 3 end clone=INA/GE-2216501 /clone_end=3' /gb=Al653487 /gl=4737466 /ug=Hs.197676 /len=306" | | |
| Cluster# | Hs.202259 | Hs.202040 | Hs.201875 | Hs.199996 | Hs.199713 | Hs.19827 | Hs.197676 | Hs.197643 | |
| Gene Name | | | | | | | | | |
| Genbank | Al700646 | Al937060 | AI825713 | AI804054 | Al921685 | W72511 | AI653487 | Al962986 | |
| Affy | 75961_at | 86612_at | 69876_at | 69600_at | 79751_at | 59623_at | 68663_at | 88622_at | |
| SegiD | 611 | 795 | 718 | 8 | 776 | 1201 | 554 | 813 | |
| #1 | 572 | 573 | 574 | 675 | 576 | 577 | 578 | 579 | |

| WO 02/059 | 271 | | 1 | 12 | | PC | T/US02/02176 |
|---|--|---|---|--|---|--|---|
| p-values 5.3291E-06 | 8.2878E-07 | 0.00678174 | 1.1628 E- 06 | 0.00380632 | 0.00223833 | 5.2381E-08 | 3.5251E-06 |
| Fold Change 0.26 | 0.08 | 4.74 | 5.05 | 0.318361618 | 0.32 | 3.13 | 0.19 |
| Cluster # Cluster Incl. A1860494.w/03b08.x1 Homo saplens 6JDNA, 3 end clonne-IMAGE-2423795 folone end=9 (\$Je3-A180494 fg]=5514100 hrefts 194274 Incl. A18457 | Hs.194083 Cluster Incl. Al968379.wu02d12.x1 Homo sapiens cDNA, 3 end /done=IMAGE-2515799 /done_end=2*/gp=45188379 /gj=5785197 incr-let 194083 Iner=577 | Hs.193745 Cluster Incl. AA4192802x05c03.s1 Homo sapiens cDNA, 3 end (done=IMAGE-756520 /done_end=9/1924A419280 /gi=2078973 /ncr=Hs.155665 lenx-456. | Hs.193602. Cluster Incl. AI761782-wi62c08.x1 Homo sapiens cDNA, 3 end clonor=IMAGE-2394830 folone_end="√">(gb=6117373 folone=IMAGE-17373 folone=I/Ne1782 folo | Hs.193491 Cluster Incl. AA701800:z334h09.s1 Homo sapiens cDNA, 3 and idone=422737 idone end=2'/gb=AA701800 /gl=2704765 increts 193401 ilen=448 | Hs.193142 Cluster Incl. AI825806:td18g01.x1 Homo sapiens cDNA, 3 and idono=INA/GE-2076048 idono=INA-GE-2076048 idono=INA-81825806 igi=5446477 idon=14s.153142 igi=707 | Hs.192872 Cluster Incl. AA719022:ah46b08.srl Homo sapiens GNNA, 3 end /donce-1292535 /done, end-e9 /gb=AA719022 /gl=2782121 /inr=Hs.192872 /len-2482 | Hs.192671 Cluster Inci. Al679896:u.69e05.xt Homo sapiens cDNA, 3 and clone=IINAGE-22555456 Idone, and-21/got=Al6778986 (gi=4899168 /ugr-Hs.192671 Aer=489' |
| Gene Name | | | | | | | |
| <u>Genbank</u> Al860484 | Al968379 | AA419260 | AI761782 | AA701600 | AI825806 | AA719022 | . Al678986 |
| Affy 69069_at | 86154_at | 67440_r_at | 88814_at | 88268_at | 85986_at | 85943_at | 68671_at |
| SeqID 741 | 817 | 108 | 651 | 202 | 719 | 211 | 587 |
| ## 280 | 581 | 582 | 583 | 584 | 285 | 586 | 587 |

| | | 110 02/032 | 2/1 | | 1 | 13 | | 10 | 1/0502/021/ |
|--|---------------------|---|---|------------|----------------------------|-----------|-------------|------------|-------------|
| | p-values | 0.00126246 | 0.00214022 | 0.00044234 | 0.00178098 | 3.57E-05 | 0.00013261 | 0.0021469 | 1.0173E-06 |
| | Fold Change | 0.309129194 | 4.256875625 | 3.32 | 3.428921588 | 3.57 | 0.212242354 | 0.23399829 | 0.25 |
| The company of the co | Cluster Description | Cluster Incl. AA778816:z;38a10.s1 Homo septiens cDDN, 3 end /donne=452538 //done_end=3'/gb=AA778816 /gl=2838147 //gehs.192714 Alen=506 | Cluster Incl. R11248;yf41c02.r1 Homo sapiens cDNA, 5 end (clone=IMAGE-129410 fictore_end=5 / gb=R1248 /gi=763983 fug+1s; 191935 flen=456° | | | | | | |
| | Cluster # | Hs.192174 | Hs.191935 | Hs,189284 | Hs.189040 | Hs.188952 | Hs.188861 | Hs.188120 | Hs.187319 |
| | Gene Name | | | | | | | | |
| | Genbank | AA778816 | R11248 | AA936632 | AI031557 | AI692624 | AA039324 | AI283643 | AI624103 |
| | Affy | 68288_at | 63131 <u>i</u> at | 68047_at | 67962 <u>g</u> at Al031557 | 79037_at | 46962_at | 87087_at | 67650_at |
| | SedID | 224 | 1102 | 260 | 320 | 298 | 2 | 410 | 535 |
| | #1 | 288 | 589 | 290 | 291 | 592 | 593 | 594 | 595 |

| | | WO 02/059 | 271 | | 114 | | | PCT/ | US02/02176 |
|---|---------------------|---|--|---|------------|------------|------------|------------|------------|
| | p-values | 1.2795E-06 | 0.00081831 | 0.00021905 | 0.02428683 | 0.00016473 | 0,00072735 | 1.4344E-05 | 3.5825E-06 |
| | Fold Change | 3.57 | 3.31 | 0.21 | 3.34 4 | 3.50 | 3.60 | 0.24 | 0.32 |
| - | Cluster Description | Octuster Incl. T79574;yd71a01.s1 Homo sapiens cDNA, 3 end kdone=IMAGE-113640 fdone_end=3 /gb=179574 /gi=698083 /ugH-8.186579 /len=533 | 3. Cluster Incl. AA420590:nc61b10.s1 Homo sapiens cDNA (clone=IMAGE:-745723 /gp=AA420590 /gi=2094496 /ug=Hs.185798 /len=501 | 3 Cluster Incl. Al381930:te72f02.x1 Homo sapiens cDNA, 3 end /done=INAGE-2092251 /done_end=3 /gb=Al381930 /gi=4194711 /ugrHs. 185708 /len=539 | | | | | |
| i | Cluster# | Hs.186579 | Hs.185798 | Hs.185708 | Hs.184343 | Hs.183918 | Hs.183412 | Hs.183409 | Hs.182809 |
| | Gene Name | | | | | | | | |
| | Genbank | T79574 | AA420590 | Al381930 | AL037594 | AA846091 | W05248 | AI653441 | A1985653 |
| | Affy | 85338_at | 78954_at | 78945_at | 92031_g_at | 90745_at | 67311_at | 85168_at | 89329_at |

SeqID

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| | | | | | 115 | | | | |
|---|---------------------|---|-------------|-------------------|-------------|-------------|-------------|------------|--------------|
| | p-values | 8,1089E-05 | 6.5741E-07 | 1.8689E-07 | 0.00011263 | 0.0000283 | 5.8955E-06 | 3.4377E-07 | 0.00108233 |
| | Fold Change | 0.22 | 0.209357681 | 0.143530024 | 3.036525383 | 0.268423966 | 0.309887046 | 3.92 | 0.214928901 |
| | Cluster Description | Hs.162364 Cluster Incl. A1083598/cox61c09.s1 Homo spopiess cDNA, 3 end idonor=INAGE;1660816 idono.end=3′fgb=A1083598 fgi=3422021 iug=Hs.298557 iden=448 | | | | | | | |
| - | Cluster # | Hs.182364 | Hs.182364 | Hs.182364 | Hs.18214 | Hs.181104 | Hs.180178 | Hs.179891 | Hs.179673 |
| | Gene Name | | | | | | | | |
| | Genbank | AI083598 | AL044670 | Al962647 | AA004622 | Al312646 | AA284268 | AI741880 | AA027103 |
| | Affy | 91173_at | 58999_at | 61317 <u>f</u> at | 59812_at | 40642_at | 54826_at | 84983_at | 63270_at |
| | SedID | 341 | 873 | 812 | ო | 124 | 95 | 626 | = |
| | #1 | 604 | 909 | 909 | 209 | 809 | 609 | 610 | 119 |
| | | | | | | | | | |

| | WO 02/0 | WO 02/059271 | | | 116 | | PC | CT/US02/02176 |
|-------------|----------------------------|--------------|---|-------------|------------|-------------|-------------|---------------|
| | p-values 1.1081E-08 | 4.4311E-05 | 0.01037591 | 0.00156659 | 0.0264336E | 6.8498E-05 | 5.1322E-05 | 0.00098285 |
| | Fold Change 3.425226104 | 0.32 | 4.628098672 | 0.325908734 | 3.33211943 | 0.268895973 | 0.326395709 | 4.67 |
| Access comp | | | Nug-Hs.1.1913. Intel-1208 Cluster Incl. Alz06063/gd/6901.x1 Homo sapiens cDNA, 3 end /done=IMAGE-1759728 Clone, end-2f /gp-7206063 /gj=3764735 | | | | | |
| i | Cluster # Hs.179222 | Hs.179153 | Hs.176067 | Hs.175048 | Hs.175044 | Hs.173975 | Hs.173369 | Hs.173088 |
| | Gene Name | | | | | | | |
| | Genbank AA292789 | N39104 | Al206063 | AI218358 | Al215667 | Al472331 | Al522299 | AA007367 |
| | Affy 62643_at | 78821_at | 92131_et | 85706_at | 86702_at | 85591_at | 92091_at | 77540_at |
| | SeqID 94 | 1073 | 388 | 383 | 391 | 483 | 499 | 4 |
| | # 612 | 613 | 614 | 615 | 616 | 617 | 618 | 619 |

| WO 02/059271 | | 1 | 17 | | PC | CT/US02/02176 |
|---|--|---|--|---|--|--|
| p-values 6.0474E-07 0.00090761 | 0.00358521 | 8.8677E-08 | 0.00028322 | 0.02233069 | 0.00013456 | 8.779E-05 |
| 5-04 Change 0.246578758 3.484499031 | 0.241109168 | 0.289343463 | 0.17 | 3.274952213 | 5.310804413 | 0.300177072 |
| Cluster Te. Cluster Description 19.17296 Cluster Ted. MRZ72(11,45641).Cxt Homo septems cDNA, 3 end febroe-IMAGE-223783 idone, and-20' igb-ABI721(11) (gj-485182, idone, and-20' igb-ABI722(11) (gj-485182, idone).Custer Ted. A44586-585677 x1 Homo septems cDNA, 3 end home-IMAGE-2145397 idone and-20' igb-A44588-58 (gj-4311437 inches 477548 homates (11) | Hs.171959 Cluster Inc. A499334-10-10008.x1 Homo sapiens cDNA, 3 end /clone=IMAGE-2178639 /clone_end=2'(pa-A499334 /gj=4391316 /nc=Hs.171859 /pm=459. | Hs.171939 Cluster Incl. A1693178.wd68411.x1 Homo sapiens cDNA, 3 end Gone=INAGE=2336757 /Gone_end=2' (pb=A1693178 /gj=4970518 /nn=Hs.171839 /qn=531" | He.171205 Custer Incl. A1038722.ox33g02.s1 Homo sapiens cDN4, 3 end /clone=IMAGE-1.638162 /clone end=2'/gp=40038722 /gj=278916 /nr=4s 171076 /an=488 | He.1709SG Cluster Incl. A1540204.td10h.12.x1 Homo sapiens cDNA, 3 end /clone=IMAGE-2075303 /clone=and=2*(9a-A1540204.tg =4457577 hcm=4*(70954.hcm26*) | Hs.170861 Cluster Incl. A4899577th01c09.x1 Homo sapiens cDN4, 3 end /done=IMAGE-2168352 /done = nd=2* (PapA1499857 /gj=4390839 /meHs.170861 /an=4.75 | Hs. (89943 Cluster Incl. AA4800092x18405.71 Homo saptiens cDNA, 5 end (clone=INA6GE-753993 forme ant=5 fgb=AA480009 fgl=2208180 Ag=Hs. (89943 Men=539" |
| Gene Name | | | | | | |
| Genbank Al672101 Al458858 | Al499334 | AI693178 | AI039722 | AI540204 | Al498957 | AA480009 |
| Affy 58916_at 83000_at | 85486_at | 58428_at | 84627_at | 92007_at | 85341_at | 47972_r_at |
| 576 576 479 | 498 | 602 | 325 | 208 | 495 | 130 |
| 620 621 | 622 | 623 | 624 | 625 | 929 | 627 |

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| | WO 02/059 | 271 | | 11 | 18 | | PC | T/US02/0217 |
|---------------------|--|------------------------|---|------------------------|---|--|---|------------------------|
| p-values | 0.00347986 | 0.00289855 | 3.6415E-07 | 1.1163E-05 | 0.00107381 | | 6.4421E-05 | 3.725341684 0.00236446 |
| Fold Change | 0.285068791 | 3.047322219 0.00289855 | 3.315735415 | 0.232010725 1.1163E-05 | 0.31 | 5.499256795 | 0.265071246 | 3.725341684 |
| Cluster Description | Cluster Incl. Al911346:wdt6a11.x1 Homo sepiens Colh4, 3 end (done=IMAGE-2328284 fdone_end=3' (gb=Al911346 /gj=5631082 /tg=18.168941 flen=396" | | Cluster Incl. Al656807.tt54b01.x1 Homo saplens cDN4, 3 end Iclone=IMAGE-2244563 Iclone_end=3'/gb=Al56807/gi=4740786 Iug=ls.167771 Ilen=531* | | Cluster Incl. Al681307.tx45g07.xrl Homo saplens ClothA, 3 end /clone=llvIAGE-2272572 /clone_end=3' /gb=Al881307 /gi=4891489 /ug=Hs.168674 /len=551' | Cluster Incl. AA424160:zv81f03.s1 Homo sapiens cDN4, 3 end /clone=IMAGE-760061 /clone_end=3 /gb=A4424160 /gi=2103148 /ug=Hs.165008 /len=447* | Cluster Incl. Al590385:ts09h02.x1 Homo saplens ClothA, 3 and /done=INAGE-2228115 /done-end=3 /gb-Al590385 /gl=4599433 /uc=Hs, 165885 /len=556 | |
| Cluster# | Hs.168941 | Hs.167899 | Hs.167771 | Hs.166784 | Hs.166674 | Hs.165909 | Hs.165885 | Hs.165805 |
| Gene Name | | | | | | | | |
| Genbank | Al911346 | A1032972 | AI656807 | AI264299 | AI681307 | AA424160 | Al590385 | AL044366 |
| Affy | 62309_at | 85068_at | 58354_at | 84903_f_at | 84314_at | 62277_at | 82441_f_at | 84763_at |

| WO 02/059 | 1271 | | | | | nc. | T/US02/021 |
|--|---|--|--|--|--|--|--|
| W G 02/03/ | 2/1 | | 1 | 19 | | | 1/0502/021 |
| <u>p-values</u> 2.4112E-05 | 0.00054065 | 1.2564E-07 | 7.3767E-05 | 0.00288569 | 0.00073954 | 0.01192547 | 0.00162269 |
| Fold Change 0.280524611 | 0.278599395 | 3.548116214 1.2564E-07 | 3.26 | 5.533470597 0.00288569 | 0.298063122 | 3.258423603 0.01192547 | 3.012339056 |
| Cluster Execution Cluster Description Fold Change He.164779 Cluster Incl. N21031;yu46106.81 Hono saplens 0.280524611 GDN A, 3 end folone-allivAE-204801 Alea He.164779 Ilea-1554 Alea He.164779 Ilea-1554 | Cluster Incl. Al039005:0x24g05.x1 Homo sapiens CDNA, 3 end /clone=IMAGE-1657304 (clone_end=3 /gb=Al039005 /gi=3278199 fureHs 164690 /len=483° | Cluster Incl. Al436670:th91b04.x1 Homo sapiens CDNA, 3 and /done=IMAGE-2125999 folone_end=3 /gb=Al436670 /gl=4283458 /ucPHs, 164369 Alen=393 | Cluster Incl. Al436297:th81c10.x1 Homo septens CDNA, 3 end /done=IMAGE-2125074 | Cluster Incl. A4610522:np93h10.s1 Homo sepiens aDNA (done=IMAGE-1133923 (pp-A4610522 /gl=2459950 /ug=Hs.162697 Ren=384 | Cluster Incl. N46855;yy73e01.s1 Homo sapiens cDNA, 3 end /donne=IMAGE-279192 //clone_end=3 /gb=N46855 /gj=1188021 //cent_s.16562 /len=524 | Hs.162130 Cluster Incl. AA5656547k25h12.s1 Homo saplens cDNA, 3 end (Jonne-INA/GE-1014599 /doing_end="3" (gb=A4656654 /gj=2337293 /uc=Hs.162130 /gn=218 | Cluster Incl. AA993596:0196910.srl Homo sapiens cDNA, 3 end /clone=IMACE-1624674 /clone_end=2/ fgb=AA993566 /gj=3180111 /ug=Hs.159983 /len=498' |
| Cluster # Hs.164779 | Hs.164680 | Hs.164369 | Hs.164226 | Hs.162697 | Hs.16262 | Hs.162130 | Hs.159983 |
| Gene Name | | | | | | | |
| Genbank N21031 | Al039005 | Al436670 | Al436297 | AA610522 | N46855 | AA565654 | AA993566 |
| Affy 62707_at | 62259_at | 84701_at | 75740_at | 84323_at | 62701_at | 84264_at | 81810_at |
| SeqID 1062 | 324 | 473 | 472 | 180 | 1078 | 168 | 268 |

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| | WO 02/059 | 271 | | | | | n.c | T/US02/02176 |
|---------------------|---|---|--|--|---|---|---|--------------|
| | WO 02/059 | 2/1 | | 1 | 20 | | rc | 1/0502/021/0 |
| p-values | 2.8623E-06 | 0.0140631 | 4.5784E-06 | 2.7534E-07 | 0.00027572 | 0.00025369 | 0.00039332 | 0.02404954 |
| Fold Change | 0.15 | 3.577532367 | 0.275099238 4.5784E-06 | 0.220904269 | 0.198202182 | 3.99 | 0.305190872 | 3.01 |
| Cluster Description | Cluster Incl. T64637:yc12h06.r1 Homo sapiens cDNA, 5 end (clone=IKAGE-80507 dione_end=5'/gb=T64637 /gi=673682 /kg=154684 /kg=154684 for=546 | Cluster Incl. Al631850:wa36h07.x1 Homo sapiens CDNA, 3 end /donn=IMAGE-2300221 /donn_e_and=3 /gb=Al631850 /gj=4683180 /trg=Hs, 156992 /en=491* | Cluster Incl. Al380583:#95g08.x1 Homo saptiens cDNA, 3 end /ctone=IMAGE-2107066 /fctone_end=3 /gb=Al380583 /gi=4190436 /lig=15.155894 /len=314* | Cluster Incl. Alg28037.wo92h01.x1 Homo sapiens CDNA, 3 end /clone=IMAGE-2462833 Holone_end=3 /gb=Alg28037 /gj=5664001 /lor+ls, 15892x flen=583 | Cluster Incl. AW026563:wv14th08.x1 Homo sapiens CDN4, 3 end /done=IMAGE-900994 flotone=IMAGE-90093 /flotone=IMAGE-90083 /flotone=IMAGE-90083 /flotone=IMAGE-90083 | Cluster Incl. Al916544:wa26h03.x1 Homo sapiens cDNA, 3 end /clone=IMAGE-2299253 | Cluster Incl. Al924465.wn56e01.x1 Homo sapiens cDNA, 3 end /clone=IMAGE-2449464 /clone_end=3' (gb=Al924465 /gi=5660429 /nreHs; 155758 /len=384' | |
| Cluster# | Hs.159367 | Hs.158992 | Hs.158984 | Hs.158832 | Hs.158741 | Hs.158549 | Hs.158258 | Hs.158113 |
| Gene Name | | | | | | | | |
| Genbank | T64637 | Al631850 | Al380583 | A1928037 | AW026553 | Al916544 | Al924465 | H15868 |
| Affy | 83908_at | 91596_at | 83727_at | 81659_at | 81648_at | 90603_at | 81591 <u>r_</u> at | 77316_at |
| SeqID | 1127 | 538 | 451 | 783 | 950 | 772 | 780 | 983 |

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| wo | 02/0 | 1592 | 71 |
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| | WO 02/0 | 059271 | | 1 | 21 | | PC | CT/US02/02176 |
|--------|---|-------------|--|------------------------|------------|-------------|-------------|---------------|
| 1 | 0.00152635 | 0.00440287 | 9.9516E-05 | 0.00216978 | 0.00692003 | 0.00015176 | 0.00013054 | 0.03882532 |
| 0 71-1 | 0.317481383 | 3.524053838 | 0.13 | 4.780974242 0.00216978 | 0.32 | 3.120542677 | 0.323852128 | 3.55 |
| | Lusser # Lusser Description -is.157437 Cluster Inc. A1367580;qv99H14x1 Homo sapiens cDN4, 3 end (John GE-1898765 /done end=3*(igh=A1367580) (gi=4137325 | | /ug=71s: 10:344 killer1409 1099 Cluster Incl. W32480/2x67603.r1 Homo sapiens cDNA, 5 end /done=IMAGE-327388 /done_ant=5/4709 /uceHs_15709 lane4709 | | | | | |
| | _ | Hs.157344 | Hs.157099 | Hs.156974 | Hs.153717 | Hs.153385 | Hs.15329 | Hs.152932 |
| | Cene Name | ٠ | | | | | | |
| 1 | Al367580 | Al367020 | W32480 | AA884688 | AA903473 | AW001002 | T83654 | Al341602 |
| | AUX 81459_at | 67164_at | 89206_at | 81383_at | 77237_at | 89891_s_at | 61965_at | 83365_at |
| 9 | 2840 440 | 439 | 1183 | 242 | 243 | 906 | 1131 | 427 |
| 7 | # 6652 | 923 | 654 | 922 | 929 | 657 | 658 | 629 |

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| WO 02/059 | 271 | | 1 | 22 | | PC | T/US02/02176 |
|--|-------------|-------------|------------|-------------|------------------------|-------------|------------------------|
| p <u>-values</u> 4.5188E-06 | 0.02012519 | 1.6074E-06 | 0.00075917 | 0.00012996 | 1.5906E-09 | 2.4725E-06 | 1.7654E-05 |
| Fold Change 5.38 | 3.248857169 | 0.237445013 | 3.80 | 3.090999056 | 0.139391318 1.5908E-09 | 0.296813014 | 0.323630776 1.7654E-05 |
| Cluster En Cluster Description Hs.152912. Cluster Inc. Cluster Description sapiens 201Na, 3 and fotone-liMAGE-208824 fotone and-9 (pp-21/10016 /pj=5108804 furnats 152912 hear-561 | | | | | | | |
| Cluster # Hs.152812 | Hs.152016 | Hs.151079 | Hs.150833 | Hs.149993 | Hs.149425 | Hs.148867 | Hs.147562 |
| Gene Name | | - | | | | | |
| Genbank Al740516 | AA603097 | R73518 | A1660245 | Al640222 | Al692813 | AI808768 | AI858012 |
| Affy 77225_at | 80845_at | 80771_at | 83226_at | 80623_s_at | 91331_at | 80487_r_at | 80322_at |
| SeqID 622 | 177 | 1117 | 999 | 543 | 009 | 889 | 736 |
| # # # | 661 | 962 | 993 | 664 | 665 | 999 | 667 |

| | WO 02/05927 | ι | | 123 | | | PCT/US | 02/02176 |
|------------------------------|---|---|---|---|--|--|---|---|
| p-values | 1.1448E-05 | 1.1834E-06 | 0.00777918 | 5.0293E-05 | 5.9089E-07 | 0.00119052 | 0.00326683 | 0.00053988 |
| Fold Change | 0.21 | 0.07 | 3.097572971 | 3.267902315 | 0.15 | 3.436009136 | 0.24 | 0.29 |
| Cluster# Cluster Description | 16.147313 Cluster Incl. A10A2492DIGFZp434G1321_r1 Horn sapters ODNA, 5 end. //danne=D/RCZpA34G1321_r1 //da>—10A2482 (gi=6421962 /ug=He.147313 //da=H682 | He.147313 Cluster Incl. AL042492:DIGFZp43461321_r1 Home sapiens cDNA, 6 and control of c | Hs.146625 Cluster Incl. A1140989;qf24h06.x1 Homo sapiens cDNA, 3 end Conne=INAGE-1751003 (done_end=v*) (gp=A1140989 (gj=S648446 liq=Hs.146525 (len=42x) | Hs.146662 Cluster Incl. A1139629:qd79g08.x1 Homo sapiens cDNA, 3 end Clone=INIAGE-175742 /clone_end="5" (gb=4/139629 (gj=5845601 /uc=Hs.14652 (len=484 | Hs.146246 Cluster Incl. A1277612:qm55ħ08.x1 Homo sapiens cDNA, 3 end Clone≔INAGE-1892703 (clone, end=v' (gb=A1277612 (gj=S899880 luc+ls: 146246 (gn=458) | Hs.146165 Cluster Incl. AI272054:qj89c08.x1 Homo sapiens cDNA, 3 end (chone≡INACE−186658 /done_end="9" (gb=dx1272054 /gj=3891221 /uc=Hs.145165 (Pen=349 | -is.146059 Cluster Incl. R83604;yp16405.r1 Homo sapiens cDNA, 5 end klong=ilklAcE+187593 klone_end=5' (gb=R83604 kgi=928481 klos=18-1461959 len=4387 | He.146042 Cluster Incl. AI080631:0x54b10.x1 Homo sapers colb.n. 3 and Achore=IMAGE-1660123 folone_and=3/ gb=AI08053 fgl=3416882 /ug=18.146042./len=482 |
| Gene Name | Ι | Ξ | I | I | Ξ. | T | I | _ |
| Genbank | AL042492 | AL042492 | Al140989 | Al139629 | Al277612 | AI272054 | R83604 | A1080631 |
| Affy | 83004_r_at | 83002 <u>ı a</u> t | 80209_at | 61978_at | 82941_at | 80134_at | 82885_at | 82879_at |
| SeqID | 298 | 867 | 366 | 364 | 408 | 404 | 1119 | 340 |
| #1 | 899 | 699 | 670 | 671 | 672 | 673 | 674 | 675 |

| | | WO 02/059 | 271 | | 12 | 24 | | PC | F/US02/02176 |
|---|--|---|---|---|--|---|---|--|--|
| | p-values | 0.00209424 | 0.01380957 | 1.7637E-06 | 6.6343E-05 | 0.04200059 | 9.1255E-05 | 0.00044029 | 0.00040144 |
| | Fold Change | 0.26 | 0.15 | 0:30 | 0.309293968 | 0.24 | 0.23 | 0.231031663 | 0.28 |
| " | Gene Name Cluster# Cluster Description | He.145989 Cluster Incl. AW0006499-w056712.x1 Homo saphers cDNA, 3 end hotone-IIMAGE-2206683 fotone_end=3' (pb=AW006499 (pj=5856277 fugr=He.145899 Iner=517 | Hs.145068 Chistler Incl. Al821472:nj04c07.x5 Homo szepiens DOIAA, s and Johnel MAGE-885366 lolone_and-3' job~482147 jgj=5440551 /ug+Hs.14508 Plen=500* | Hs.144871 Cluster Incl. Al686114:182c02.x1 Homo saplens cDNA, 3 end kotone-IIMAGE:2248994 kotone-end-3' kgb-Al686114 kjg-4897408 kug-Hs.144871 llen-4897 | Hs.14864 Cluster Incl. ANV052142xw28d08.x1 Homo sapiens DDNA, 3 and vloine=NIAGE-2544783 /done_and=?/p=AW052142 (gj=5914501 /ug=Hs.144864 /len=549* | Hs.144151 Cluster Incl. AI868620.yo63n06.x6 Homo sapiens cDNA, 3 erd /done=IMAGE-181891 /done=and=3 /gb=4088620 /gj=4827928 /ugeHs.144151 RenE817 | Hs.143995 Cluster Incl. AI570222:t076e03.x1 Homo sapiens cDNA, 3 erd Iodone=IIMAGE-2:184220 Iodone_end=3' (pd-AI570222 gj=4633596 IouzHs. 143965 fen=458 | Hs.143973 Cluster Incl. AI740621 xug23e12.x1 Homo sapiens DINA, 3 end vlomor=IIIAAGEZ385965 idene_and=97 igp=AI740621 ig=5108909 ing=Hs.143873 ilen=4597 | Hs.143789 Cluster Incl. AB94068-WG7Cf0.x1 Homo saplens of ADA, 3 and Idone=IMAGE-22386588 Idone and=7 (pp-44694059 Igi=4971399 lug=Hs.143789 Nen=540* |
| | Gene | | | | | | | | |
| | Genbank | AW006499 | AI821472 | AI686114 | AW052142 | AI668620 | AI570222 | AI740621 | AI694059 |
| | Affy | 82860_at | 89087_at | 77106_at | 79803_at | 77077_at | 82595_at | 79618_at | 77053_at |
| | SeqID | 918 | 712 | 593 | 954 | 571 | 619 | 623 | 604 |
| | #1 | 929 | 677 | 678 | 629 | 089 | 189 | 682 | 683 |
| | | , | • | | | | | | |

| | | | | | 125 | | | | |
|---|---------------------|--|---|--|-------------|-----------|------------|------------------------------|-------------|
| | p-values | 9.8109E-05 | 4.2869E-07 | 0.00038894 | 0.00182392 | 5.382E-07 | 0.00918651 | 0.0002078 | 0.00024859 |
| | Fold Change | 6.44 | 0.22 | 3.543032267 | 0.326197997 | 0.29 | 0.33 | 0.29343002 | 3.063096074 |
| | Cluster Description | Cluster Incl. AA633203:nq57b02.s1 Homo saptens colbA, 3 end /clone=IMAGE-147947 /clone_end-3 /gb-AA633203 /gi=2556617 /ug=Hs.14258 llen=570° | clusier inci. AL040178:DKFZp434F0213_s1 Homo sapiens cDN4, 3 end chone=DKFZp434F0213 cione_end=3" cbo=AL040178 gi=5403143 lug=Hs.142003 le=738 | Cluster Incl. Al963725.wr66b11.x1 Homo expiens cDNA, 3 end idone=IMAGE-2492637 Iclone_end=3' igb=Al963725 igi=5756438 iugh+s_207705 ilen=427 | | | | | |
| - | Cluster# | Hs.14258 | Hs.142003 | Hs.141024 | Hs.140996 | Hs.13821 | Hs.137907 | Hs.13766 | Hs.137447 |
| | Gene Name | | | | | | | | |
| | Genbank | AA633203 | AL040178 | Al963725 | AA412205 | W63684 | Al018237 | AA015613 | Al539443 |
| | Affy | 90421_at | 90389_at | 69928_at | 75277_f_at | 91422_at | 75195_at | 65185 <u>_g_</u> at_AA015613 | 61191_at |
| | SeqID | 193 | 864 | 814 | 103 | 1192 | 316 | ဖ | 507 |
| | # | 684 | 685 | 989 | 289 | 989 | 689 | 069 | 691 |

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|----|----|-----|----|----|

| | WO 02/059 | 271 | | 1 | 26 | | PC | T/US02/02176 |
|------------------------------|--|--|---|---|--|--|--|---|
| b-values | 1.3855E-05 | 0.00011435 | 0.00068507 | 0.00145442 | 0.00038716 | 0.02123852 | 0.03600705 | 8.8297E-05 |
| Fold Change | 0.31404932 | 0.25 | 0.24 | 5.00 | 0.127900019 | 3.05 | 4.015935457 | 0.260125725 |
| Cluster# Cluster Description | Hs.137262 Cluster Ind. N49691;yy58d01.s1 Homo sapiens cDNA, 3 end /done=INAGE-277729 /done_end=3 /gb=H49691 /gj=1190757 /ug=Hs.197782 /en=4689 | Hs.137003 Cluster Incl. AI242023:qh81a04.x1 Homo sapiens cDNA, 3 and /donne=INAGE-1851054 /done_end=3 /gb=AI242023 /gl=3837420 /ug=Hs.137003 /en=A48 | Hs.135657 Cluster Incl. AAG34591;nf81b04.s1 Homo saplens cDNA, 3 and /dons=IMAGE-926287 /done_end=3 /gb=AAG34591 /gj=2278844 /ug=Hs.135697 /len=A929. | Hs.13561 Cluster Incl. Al819340.wg61e01.x1 Homo saplens cDNA, 3 end /donne=IIAAGE-2369544 /clone_end=3 /gb=Al819340 /gj=5438419 /ug=Hs.13561 /len=559 | Hs.135405 Cluster Incl. AI057450.cw80cd3.x1 Homo saplens cDNA, 3 end /clone=IIAAGE-1653124 /clone_end=3 /gb=AI057450 /gl=3331316 /ucrl+s.136406 /len=379 | Hs.135056 Cluster Incl. H063503y179g02.r1 Homo sapiens CDNA, 5 end (clone=IMAGE-44306 /clone=Ind=7 /gb=+H06350 /gj=869902 /dp=18.7 /gb=+H06350 /gj=869902 /dp=18.7 /gb=H06350 /gj=869902 | Hs.134665 Cluster Incl. AI673818:to73f12.x1 Homo sapiens cDNA, 3 end /clone=IMAGE-2183951 /clone_end=3 /gb=AI673818 /gj=4853549 /wz=Hs.134565 /en=2017 | Hs.134110 Cluster Ind. AUT/36121:co224d01.xr Homo sepiers cDVIA,3 end Idone=IMAGE-1676257 Idone end-2* Igb=M01211 Igi=3412529 Aug=Hs.134110 Aen=457 |
| Gene Name | | | | | | | | |
| Genbank | N49591 | Al242023 | AA534591 | AI819340 | Al057450 | H06350 | Al673818 | Al078121 |
| Affy | 79133_at | 82436_at | 82385_at | 78442_at | 78844_at | 86587_at | 78668_r_at | 78555_at |
| SeqD | 1080 | 396 | 157 | 708 | 331 | 979 | 284 | 337 |
| # | 692 | 693 | 694 | 695 | 989 | 269 | 869 | 669 |

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|----|-----|-----|-----|

| WO 02/059 | 271 | | 1 | 27 | | PC | T/US02/02176 |
|--|-----------|------------|------------|---------------------|-------------|-------------|--------------|
| 0.00810637 | 1.664E-06 | 0.00198223 | 4.0319E-09 | 0.00197051 | 0.03333431 | 0.00161436 | 0.00059354 |
| Fold Change 4.039586551 | 0.07 | 3.87 | 10.36 | 3.15 | 0,305626072 | 0.280881931 | 3.140512538 |
| Cluster # Cluster Description Hs.134053 Cluster Incl. AI023295.ow95b12.xrl Homo sapleins CDNA, 3 and kone=INAGE-1654559 Idone_and=3 (\$p=\000000000000000000000000000000000000 | | | | 0 0 | | | |
| Cluster# Hs.134053 | Hs.133471 | Hs.13337 | Hs.133294 | Hs.133294 | Hs.133226 | Hs.13299 | Hs.132586 |
| Gene Name | | | | | | | |
| Genbank Al023295 | AI758223 | AI276259 | Al053741 | Al690773 | Al499220 | H92987 | AI031771 |
| Affy 78538_at | 76769_at | 90224_at | 91875_s_at | 91873_s_at Al690773 | 78391_at | 65173_at | 78231_at |
| SeqID 318 | 643 | 407 | 329 | 296 | 496 | 966 | 321 |
| 4 200 | 701 | 702 | 703 | 704 | 705 | 708 | 707 |

| | | | | 12 | 28 | | | |
|-----------------------|---|---|--|--|--|------------|---|--|
| p-values | 6.3592E-07 | 0.00126632 | 0.00124922 | | 2.9214E-05 | 0.00029449 | 4.2586E-09 | 0.01303714 |
| Fold Change | 0.16 | 3.09 | 0.271854365 0.00124922 | 0.30 | 0.080787812 | 7.28 | 0.103783146 | 3.015818659 |
| # Cluster Description | 87 Cluster Incl. Al890418:wm85ar1z.x1 Homo sapiers cDNA, 3 end Iclone=IMAGE-2442718 Iclone_end=3' (40=8890418 /gl=589585z /ug=Hs: 131887 /len=516" | 86 Cluster Incl. AW003102:wr03e03.x1 Homo saplens cDNA, 3 end /clone=IMAGE-2480476 /clone_end=3' (49=4W003102 (gi=5849940 /ug=Hs, 131886 /en=612. | 70 Cluster Incl. Al937390:wp76f01.x1 Homo saphers cDNA, 3 end Idone=IMAGE:2467705 Iclone_end=3' Id92Al937390 /gl=5676260 /ug=Hs.131170 Ilen-483' | 44 Cluster Incl. Al631301:tz82f05.x1 Homo sapiers cDINA, 3 end (done=IMAGE-2295105 /done_end=3'/qb=46831301 /gi=4682631 /ugeHs.131044 /len=462' | 893 Cluster Incl. A1124631:am59f08.xr Homo sapiens cDNA, 3 end /done=IMAGE=1539879 /done_end=3 /do=A1124631 /gi=3593145 //geHs.13083 /len≠416 | | 999 Cluster Incl. AA621478:af92e12.s1 Homo saphens cDNA, 3 end /done=IMAGE-1055278 /done_end=3'\tilde{g}=2525417 /uge+hs.130699 len=398* | 316 Cluster Incl. AI572165:te37d05.x1 Homo sapiers cDNA, a end fatne=IMAGE-2088873 Idone_end=3'40p-AI572166 (gi=4535530 /ug=Hs.130316 .len=34* |
| Cluster# | Hs.131987 | Hs.131886 | Hs.131170 | Hs.131044 | Hs.130893 | Hs. 130853 | Hs.130699 | Hs.130316 |
| Gene Name | | | | | | | | |
| Genbank | AI890418 | AW003102 | Al937390 | AI631301 | Al124631 | AI810266 | AA621478 | AI572156 |
| Affy | 76703_at | 74698_at | 77926_at | 82120_at | 59911_f_at | 82094_i_at | 47481_at | 90691_at |
| SeqID | 763 | 606 | 797 | 537 | 353 | 693 | 2 8 | 521 |
| #1 | 708 | 500 | 710 | 711 | 712 | 713 | 417 | 715 |

| WA | 02/05 | 027 |
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| | WO 02/059 | 271 | | 12 | 29 | | PC | T/US02/02176 |
|---------------------|--|-------------|--|------------|------------|--------------------|-------------|--------------|
| p-values | 0.00155658 | 0.00247835 | 0.00011441 | 7,7763E-10 | 1,9965E-06 | 2.399 E- 05 | 5.2644E-08 | 0.00037957 |
| Fold Change | 3.808419817 | 4.615880703 | 0.29 | 10.43 | 0.25 | 4.421237061 | 0.054094404 | 3.09 |
| Cluster Description | Hs.130123 Cituster Incl. AA906874:;0j82c12.s1 Homo sapiens cJNN, 3 end Idonne=IMAGE-1504822 /done_end=3'/gp=A4906874 /gi=3040997 /ug=Hs.130123 /len=3689 | | Cluster Ind. Al492154;tg12f12.x1 Homo sapiens cDN4, 3 end (clone=INAGE-2108591 fclone_end=3'/g0=Al492164/gi=4393157 /uq=hs;128327 Men=547 | | | | | |
| Cluster# | Hs.130123 | Hs.129873 | Hs.129327 | Hs.128022 | Hs.127630 | Hs.127307 | Hs.127229 | Hs.126798 |
| Gene Name | | | | | | | | |
| Genbank | AA905874 | Al820661 | Al492154 | AI859620 | AA928776 | Al263819 | Al300876 | AA975530 |
| Affiy | 77715_at | 61908_at | 74462_at | 82009_at | 91844_at | 90541_at | 77135_at | 74051_at |
| SedID | 245 | 709 | 491 | 739 | 258 | 400 | 417 | 265 |
| # | 716 | 717 | 718 | 719 | 720 | 721 | 722 | 723 |

| | | WO 02/059 | 271 | | 13 | 10 | | PC | T/US02/02176 |
|---|---------------------|--|--|--|---|---|---|--|---|
| | p-values | 1.2217E-06 | 2.3169E-05 | 0.00040139 | 7.0947E-06 | 4.2446E-06 | 4.7816E-06 | 1.6889E-06 | 4.0015E-05 |
| | Fold Change | 0.19 | 3.31847909 | 5.050317981 | 3.297301166 | 0.26 | 0.169055931 | 0.25 | 0.32 |
| | Cluster Description | Cluster Incl. AI599858.tm8 1b11.x1 Homo sapiens GDNA, 3 end /clone=IMAGE-2164509 /clone_end=3 /gb=AI599858 /gj=4598906 /ug=Hs; 126768 /len=4597 | Cluster Incl. AA913703:ol38e01.s1 Homo sapiens GDNs, 3 end /done=IMAGE-1525752 /clone_end=3 /gb=AA913703 /gj=3053095 /tg=Hs.126733 /lan=507* | Cluster Incl. AI792817:ol64f01.y5 Homo saptiens CDNA, 5 end /donë=IMAGE-1534393 /clone_end=5 /gb=AI792817 /gi=5340533 /kg=Hs,126672 Aen=306 | Cluster Incl. AA905481:ok01h09.s1 Homo sapiens 6.Dbh, 3 end /done=IMAGE-1506593 Actone_end=3' /gb=AA905481 /gi=3040604 /ug-Hs, 126390 /lan-461' | Cluster Incl. Al934342:wp04e12.x1 Homo sapiens cDNA, 3 end /donne=IMAGE-2463886 /donne_end=3 /gb=Al934342 /gi=5673212 /nc=1s, 128780 /len=521* | Cluster Incl. A4584403·nn81a05.s1 Homo sapiens cDNA, 3 end /clone=IMAGE-1090256 /clone_end=3' /gb=A4584403 /gi=2369012 /ug=Hs.125376 /len=512' | Cluster Incl. AA844007.a81409.s7 Homo septens cDNA, 3 end /donne-IMAGE-1388177 /clone_end=3' /gb=AA844007 /gi=2930458 /tig=hs; 124438 flen=310° | Cluster Incl. AA848010:od24g07.s1 Homo sapiens cDNA (clone=IMAGE-1388924 /gb=AA848010 /gj=2834528 /ug=Hs.124250 /fen=415 |
| ! | Cluster# | Hs.126768 (| Hs.126733 (| Hs.126672 (| Hs.126390 | Hs.125780 | Hs.125376 | Hs.124436 | Hs.124250 |
| | Gene Name | | | | | | | | |
| | Genbank | AI589858 | AA913703 | AI792817 | AA905481 | Al934342 | AA584403 | AA844007 | AA848010 |
| | Affy | 90069_at | 46538_at | 77013_at | 90479_at | 73933_at | 76770_at | 73801_at | 73794_at |
| | SedID | 526 | 252 | 699 | 244 | 786 | 172 | 238 | 240 |

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| wn | 02 | 0.5 | 02 | 71 |
|----|----|-----|----|----|

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|------------------------|-------------|-------------|------------|------------|------------|------------|------------|
| p-values 4.4423E-07 | 1.5522E-06 | 4.2459E-08 | 1.9577E-06 | 2.1736E-10 | 0.00094679 | 7.1425E-07 | 0.00166257 |
| Fold Change 0.09 | 5,302996355 | 0.100992647 | 0.21246828 | 0.17 | 3.60100109 | 0.24 | 0.29523189 |
| Cluster # | | | | | | | |
| Cluster# Hs.124141 | Hs.124015 | Hs.123933 | Hs.12369 | Hs.122593 | Hs.121532 | Hs.121518 | Hs.120959 |
| Gene Name | | | | | | | |
| Genbank Al458003 | Al953838 | Al809953 | AA707308 | AL037998 | AA033764 | H41870 | Al278074 |
| Affy 90009_at | 59471_at | 57605_at | 65155_at | 81670_at | 90316_at | 81589_at | 74760_s_at |
| SeqID 477 | 807 | 691 | 509 | 828 | 16 | 686 | 409 |
| 732 | 733 | 734 | 735 | 736 | 737 | 738 | 739 |

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| | | WO 02/0592 | 271 | | 13 | 2 | PCT/US02/02176 | | |
|-----------|--------------------------|--|--|--|--|---|--|--|--|
| | p-values | 3.0941E-06 | 1.5413E-05 | 2.9798E-06 | | 4.3863E-08 | 0.00159704 | 1.3663E-05 | 1.5912E-05 |
| | Fold Change | 0.22 | 0.285003866 | 0.239883875 | 0.24 | 0.198179833 | 3.768700804 | 0.204140593 | 0.240592502 |
| 1 1 1 1 1 | ter# Cluster Description | Hs.12082 Cluster Incl. AI201982;qs79e02.x1 Homo spepiens cDNA, 3 end idone=IMAGE;1944314 idone=ImAGE;1944314 idone=Ind=3/(gb=AI201982/gj=3754588 idg=Hs.12318 incm=480 | Hs.120785 Cluster Incl. W73890.zd65h02.s1 Homo sapiens cDN4, 3 end idane=INAAGE-345555 Idane_end=3' (gb=W73890 /gj=1382285 iug=Hs.120785 /len=570 | Hs.120568 Cluster Incl. Al302387.cpt50g05.x1 Homo sapiens cDNA, 3 end (donne=INAGE+1901720 /clone_end=3'/gp=43102387/gj=3961733 /ug=Hs.120568 /len=396 | Hs.120388 Cluster Incl. AI742521;wg43h08.x1 Homo sapleins 6DNA, 3 and idone=il/AGE-2387903 idone_ent=3 /gb=AI742521 /gi=5110809 /ug=Hs.120388 /len=549 | Hs.12024 Cluster Ind. R42914;yg14a10.s1 Homo sepiens cDNA, 3 end klonne=IMAGE-32117 klonne_end=3' (gb=R42914 /gj=819821 klg=Hs.12024 flen=514* | Hs.118599 Cluster Incl. Al821005;yb594083y5 Homo sapiens cDNA, 5 end /donne=IIAAGE-75477 /done_end=5'/gb=Al821005 /gl=5440084 /dg=Hs.118599 /nen=507 | Hs.118513 Cluster Incl. A1161387.qb80c11.x1 Homo septiens cDNA, 3 and idono=IIIAAGE-1708420 idono_ent=3'/gb=A1161387/gj=8894611 /ug=Hs.118613 /len=693' | Hs. 118502. Cluster Incl., NuoCa21883-wg78-d5.x1 Homo sapiens cDNA, 3 end /done=IMAGE-2555504 /done_end=2/ (pi=AW0C2186 /gi=S914545 /ug=Hs. 118502 /len=485 |
| | Cluster# | Hs.12 | Hs.12(| Hs.120 | Hs.120 | Hs.12 | Hs.11 | Hs.11 | Hs.11 |
| | Gene Name | | | | | | | | |
| | Genbank | Al201982 | W73890 | Al302387 | AI742521 | R42914 | AI821005 | Al161367 | AW052186 |
| | Affy | 76326_at | 57550_at | 66390_at | 88669_at | 61879_at | 90251_at | 52946_at | 76076_at |
| | SedID | 387 | 1207 | 419 | 632 | 1107 | 710 | 377 | 955 |
| | #1 | 740 | 741 | 742 | 743 | 744 | 745 | 746 | 747 |

| | WO 02/0592 | 271 | | 13 | 3 | | PC | Γ/US02/0217 |
|-----------------------------------|---|--|---|--|---|---|---|--|
| p-values | 3.2934E-06 | 3.2676E-06 | 0.00404612 | 0.00013207 | 0.00138841 | 0.00090085 | 0.00320729 | 1.5411E-07 |
| Fold Change | 0.223181865 | 3,141681584 | 0.314306045 | 0.305498776 0.00013207 | 0.325353873 | 0.26 | 0.322106515 0.00320729 | 0.090939502 |
| Cluster # Cluster Description | Hs.118392. Cluster Incl. Al824037-wi29th0z.x1 Homo sapiens cDNA, 3 end clone=INAGE-2404275 /clone_end=3' (gb~4824037 /gj=5444708 /ug=Hs.118392 /len=603 | Hs.118338 Cluster Incl. NZ4987.yx16d11.17 Homo sapiens 3.141681584 cDNA, 5 end Cichon=IMAGE-261909 Clone_end=5" (pl=MZ4987 /gj=1139137 /ug=Hs.118338 /lap=887 | Hs.118262 Cluster Incl. Al698243:wa70g05.x1 Homo sppiens cDNA, 3 end ckonce-IN/AGE-2301560 /clone_end=v? (gb=46986243 /gi=4986143 /ug=Hs.118202 /len=431* | He.118121 Cluster Incl. A1124882:am57708.x1 Homo sapleans cDNA, 3 and IdonperIMAGE-1538687 Idone_end=3' (gb=A1124882 (gl=5583396 Mg=He.118171 Nen=408 | Hs.11782 Cluster Incl. Al823572×wh55g06.x1 Homo sappiens cDNA, 3 end clonne=INAGE-2384698 clone_end=9**(\$p→s1823572 /gj=5444243 lug=Hs.11787 ilen=538* | Hs.117687 Cluster Incl. AI766029.wh67b04.x1 Homo sapiens clNN ₄ , 3 end clonperII(AIGE-2388775 idone_end=3' (gb=AI766029 (gl=S232538 iug=Hs.117887 len=704 | Ha.117474 Cluster Incl. R49146;yg69h06.s1 Homo sapiess cDNA, 3 and clonne=INA/GE-38818 /clone_end=3'/gb=R49146 /gj=820214 /ug=Hs.117474 /len=439 | Hs.116123 Gluster Incl. AI016755:ov27006.x1 Homo sapiens cDNA, 3 end /donne=INAGE-1638538 /done_end=3'/gb=A016755 /gj=323.1091 /ug=Hs.116123 /len=4/77 |
| Gene Name Cl | 圣 | £ | 至 | 至 | Ĭ | ¥ | ¥ | 至 |
| Genbank | AI824037 | N24987 | AI698243 | Al124882 ° | AI823572 | AI766029 | R49146 | Al016755 |
| Affy | 57528_at | 47434_at | 76029_at | 76015_at | 57517_at | 76163_at | 66305_at | 66240_s_at |
| SeqID | 716 | 1066 | 609 | 354 | 713 | 922 | 1108 | 315. |
| ## | 748 | 749 | 750 | 751 | 752 | 753 | 754 | 755 |
| | | | | | | | | |

| | 110 02/00/ | | | 13 | 34 | | | 1.0002,021.0 |
|---------------------|--|--|---------------------|------------|------------|------------|------------|--------------|
| p-values | 7.063E-09 | 2.0933E-09 | 0.00377036 | 6.3947E-05 | 5.4104E-05 | 5.4138E-06 | 0.00036834 | 0.00069299 |
| Fold Change | 0.139924253 | 8.54 | 4.52 | 0.21 | 0:30 | 0.25 | 0.18 | 0.262512195 |
| Cluster Description | Hs.116110 Cluster Incl. Al916626.wa28h02.x1 Homo sapiens cDNA, 3 end /cione=INAGE-Z289443 /cione_end=3'/gp=4819626 /gj=5636481 /ug=Hs.116110 /en=487 | Cluster Ind. Al275140:q/70h02.x1 Homo saplers cDNA, 3 end /done=IMAGE-1877715 idone_end=3 /gb-Al275140 /gj=5897414 /ug=16.116104 hen-444* | | | | | | |
| Cluster# | Hs.116110 | Hs.116104 | Hs.115838 | Hs.115497 | Hs.115315 | Hs.115173 | Hs.114889 | Hs.11455 |
| Gene Name | | | | | | | | |
| Genbank | Al916626 | AI275140 | Al344053 | Al498375 | AI978710 | Al418405 | AI797063 | T64447 |
| Affy | 90168_at | 88580_at | 88567_s_at Al344053 | 76118_at | 81061_at | 89807_at | 81039_at | 54983_at |
| SeqID | 773 | 405 | 429 | 494 | 832 | 464 | 673 | 1126 |
| # | 756 | 757 | 758 | 759 | 760 | 761 | 762 | 763 |

| | | | | | 13 | 55 | | | |
|--|---------------------------|---|--|------------|---|--|-------------|---|-------------|
| | p-values | 1.6712E-08 | 5.1114E-07 | 1.9391E-08 | 2.882E-07 | 6.2396E-05 | 0.00068815 | 2.0122E-05 | 0.00718188 |
| | Fold Change | 0.093652816 | 0.200282848 | 0.20 | 0.202348411 | 0.084843922 | 3.166594901 | 0.116868505 | 0.269236792 |
| many or and the state of the st | · · · Cluster Description | Hs.113750 Cluster Incl. Al091164;oo18a08.x1 Homo sapiers cJNA, 3 end /donne=INAAGE-1686518 /done_end=3/gp=Al091164/gi=3430213 /ug=Hs.113790 /lone=437 | Hs.112895 Cluster Incl. AI742490.wg43e10.x1 Homo sapiens cDNA, 3 end /dome=INAGE-2367882 /done_end=3′/gb=AI742490 /gi=5110778 /ug=Hs.112895 /len=550 | | Cluster Ind. AA452295;zx30c10.r1 Homo sapiens cDNA, 5 end /done=INA/GE-787986 /done_end=5'/gb=AA452295 /gi=2165964 /ug=Hs.110406 /len=528 | Cluster Incl. Al732274;yj88b10.x5 Homo saplens CoDNA, 3 end Iclone=INAQE-153883 Iclone_end=3' /gb=Al732Z14 /gi=5053387 /ug=Hs, 11006 Alen=538" | | i Cluster Incl. Al871044:wI78h03.x1 Homo saptens cDNA, 3 end /clone=IMAGE-2431061 /clone_end-3 /gb=Al871044 /gl=5545012 /ug+Hs.109525 /len=486 | |
| | Cluster # | Hs.113750 | Hs.112885 | Hs.112572 | Hs.110406 | Hs.11006 | Hs.109653 | Hs.109525 | Hs.107253 |
| | Gene Name | | | | | • | | | |
| | Genbank | AI091154 | AI742490 | AA628467 | AA452295 | AI732274 | N91175 | AI871044 | AA059401 |
| | Affy | 75585_at | 57022_at | 80917_at | 56941_at | 48115_at | 42353_at | 56910_at | 63344_at |
| | SedID | 345 | 631 | 189 | 124 | 818 | 1098 | 750 | 32 |
| | ## | 764 | 765 | 766 | 767 | 168 | 492 | 022 | 134 |

| WO 02 | /059271 | | | 136 | | | PCT/US02/02176 | |
|-------|---------|-------|-------|-------|-------|-------|----------------|--|
| 33978 | 7E-05 | 7E-05 | 2E-06 | 26278 | 1E-05 | 5E-05 | 1E-05 | |

| | | | | 1. | 90 | | | |
|---------------------|---|---|--|--|------------|-------------|-------------|------------|
| p-values | 0.00333978 | 3.0927E-05 | 3.2707E-05 | 2.4582E-06 | 0.04026278 | 6.4561E-05 | 4.2545E-05 | 7.7541E-05 |
| Fold Change | 0.289956429 | 0.212581687 | 0.25026049 | 3.113529847 | 0.31991901 | 0.194470029 | 0.147540619 | 0.29 |
| Cluster Description | Cluster Ind. AA173572::;p04e02.s1 Homo sapiens cDN4, 3 and /ohone=IMAGE-595418 idone_end=3'/gb=AA173572 /gi=1753704 /ug=16.10633 /fen=595 | Cluster Incl. AA806965:0c34c06.s1 Homo sapiens cDN4/ (clone=IMA/GE-1351584/gp=AA806965 /gi=2876541 /ug=Hs.106771 flen=569 | Hs.103395 Cluster Incl. AA147751;zJ50f03;s1 Homo saplens cDNA, 3 end Clone=IMAGE-505373 /done_end=3' (gb=AA147751 gl=1717314 /ug=Hs.103395 len=538 | Cluster Incl. Al885164:w/80e04.x/I Homo sapiens cDNA, 3 end /clone=IMAGE-2432190 /clone_end=3 /gb=Al885164 /gi=5590328 /kg=Hs.103305 /len=506 | | | | |
| Cluster# | Hs.10683 | Hs.106771 | Hs.103395 | Hs.103305 | Hs.102793 | Hs.102541 | Hs.102367 | Hs.10198 |
| Gene Name | | ٠ | | | | | | |
| Genbank | AA173572 | AA806965 | AA147751 | Al885164 | AI707589 | AI752682 | W72347 | Al201965 |
| Aff | 48063_at | 58174_at | 48040_at | 56190_at | 58429_at | 63315_at | 48032_at | 87970_at |
| SeqID | 75 | 529 | . 62 | 753 | 615 | 642 | 1199 | 386 |
| # | 772 | 773 | 774 | 775 | 776 | 111 | 778 | 779 |

| | WO 02/059 | 271 | | 1 | 37 | | PC | T/US02/02176 |
|-------------------------------|---|---|---|---|---|--|--|--|
| p-values | 6.125E-07 | 2.9689E-05 | 1.6462E-05 | 0.00415608 | 0.00016062 | 4.0981E-05 | 0.00451133 | 0.00170328 |
| Fold Change | 0.244166495 | 0.275558723 | 13.54323165 | 8.54 | 8.025997484 | 5.726391287 | 3.734521363 | 3.65368217 |
| Cluster # Cluster Description | Cluster Incl. / sapiens cDN /clone_end= | Hs.101889 Cluster Incl. A879723:to41b10.x1 Homo sapiens cDNA, 3 end /done=IMAGE-2067163 /done=IMAGE-2067163 /clone end-3* (JehA379723 /gi=4189576 /clone and-3* (JehA379723 /gi=4189576 /clone and-3* (JehA379723 /gi=4189576 /clone and 3* (JehA379723 /gi=4189576 | Applants, for the property of | Cluster Inc. AA687785es7207.srl Homo sapiens cDNA, 3 end kdone=IIMAGE-969733 Kolone, end-97197777777777777777777777777777777777 | Appl. 1923 100 100 100 100 100 100 100 100 100 10 | Cluster Inc. AB29169.wk78b02.x1 Homo sapiens cDNA, 3 end /done=IMAGE-2421291 /done=IMAGE-2421291 /done=IMAGE-24201291 /done=id=2798040 /do=6249840 /do=6249840 /do=6249840 /do=6249840 /do=6249840 | Cluster Inc., AA/01/1882/380804.s1 Homo sepiens cDNA, 3 end /clone=461/66 /clone=461/61 end end=3/gheAA/01/188 /gj=2704353 /cl-16.400977 Inc210. | May 18, 1900 May 1900 |

AA663786

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Gene Name

Genbank AA418636

Affy 63313_at

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| WO 02/059 | 271 | | 138 | | | PCT/U | PCT/US02/021 | | | | | |
|--|--|---|---|--|--|--|--|--|--|--|--|--|
| p-values 8.1229E-05 | 0.00070373 | 9.1295E-07 | 1.7708E-06 | 3.3328E-06 | 7.1154E-05 | 1.3678E-06 | 0.182339389 5.8223E-06 | | | | | |
| <u>Fold Change</u> 3.149965078 | 0.32 | 0.317111476 | 0.287990459 1.7708E-06 | 0.264521584 | 0.233645562 7.1154E-05 | 0.185193232 | 0.182339389 | | | | | |
| Cluster # — Cluster Description Cluster Incl. Al025 (95:xv40f01 xr1 Homo sapeins DJN, 3 end clone=MAAGE-1639801 clone_end=9' (95-Al02519) (gj=52.40812 And-Hs. 31 604, flent=227 | Cluster Incl. AA081045;zn33e02.s1 Homo sapiens cDNA, 3 end /donn=IMAGE:468242 /donne_ente-2/pen-A081045/gj=1823033 /irr=He 149294 Apr=45081 | Cluster Ind. W44859.2x239809.r1 Homo sapiens cDNA, 5 end /clone=IMAGE-323704 /clone=IMAGE-323704 /clone=IMAGE-323704 /clone=IMAGE-323704 /clone=IMAGE-323704 /clone=IMAGE-323704 /clone=IMAGE-323704 /clone=IMAGE-3230706 | Mayers, House 2008 DKF2p434C1123_11 Cluster Incl., ALDAS086DKF2p434C1123_11 Homo sapates 2DMA, Ed. Idones—DKF2p434C1123 Idones_ender5 Idones—DKF2p434C1123 Idones_ender5 Idones_EDKF2p436C1123 Idones_EDKF2p436C1123 Idones_ender5 Idones_EDKF2p436C1123 Idones_EDKF2p436C1123 Idones_EDKF2p436C | Cluster Incl. W56090.zc58g03.r1 Homo saplens cDNN, 5 end /done=INA/GE-228356 /done_ind=0.pd=7/gb=W56090 /gi=1357999 Air=Hs 17310 /len=8201 | Cluster Incl. AA007390:zh99a06.r1 Homo sapiens CDINA, 5 end clonne=INAGE-429394 /done_ind=2'/gb=A4007390 /gj=1493628 /m=14-4 -1650.44 /lan=552 | Cluster Inc. AA4303 14.2w68h06.r.1 Homo Cluster Inc. AA4303 14.2w68h06.r.1 Homo sapiens cDNA, 5 end (clone=IMAGE-781403 Idone pin=2 (pge=AA430314 (gi=2113524 incette 69796 fine=579 | Cluster Incl. W07304:za97b10.r1 Homo sapiens cDNA, 5 end /clone=IMAGE-300475 | | | | | |
| Gene Name | | | | | | | | | | | | |
| Genbank Al025199 | AA081045 | W44656 | AL043089 | W56090 | AA007390 | AA430314 | W07304 | | | | | |
| Affy 66998_at | 79496_r_at AA081045 | 47889_at | 65867_at | 57586_at | 82582_at | 48927_at | 53793_at | | | | | |

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| p-values | 6.1089E-05 | 2.7057E-06 | |
|--|--|---|--|
| Fold Change p-values | 0.14 | 0.110161171 | |
| Gene Name Cluster# Cluster Description | Cluster Incl. N50065;yz10h03.s1 Homo sapiens cDNA, 3 end /clone=IMAGE-282677 | Agnite, 169722 Mer-1860 Agnite, 169722 Mer-1860 Cluster Inch. Ty CZ255/9/1608 st Homosapiens 0.110161171 2.7067E-06 GDNA, 3 end Actone-ill-MAGE-156771 | /clone_end=3'/gb=R70255 /gj=843772 /ug=Hs.25150 /len=367' |
| Gene Name | | | |
| Genbank | N50065 | R70256 | |
| Affy | 84535_at | 52449_at | |
| SedID | 1081 | 1116 | |
| # | 962 | 797 | |

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| p-value | 5.020000E-05 | 3.510000E-07 | 2.660000E-08 | 2 3800001-05 | | 7.124464E-06 | | 1.340000E-05 | | 6.210594E-06 | | 4.710000E-08 | 1.380000E-06 | 4.050474E-06 | 7.807645E-06 | 4.150000E-04 | 1.318202E-04 | • | 4.120000E-07 | 3.247311E-05 | 1.610000E-04 | 1.518608E-06 | 3.900000E-07 | 2.613853E-08 | 7.205058E-06 | 1.770803E-06 | 4.329042E-05 | | 1.000000E-04 | 4.436191E-07 | 2.373819E-06 | 1.093627E-04 | 1.086072E-05 |
|--|------------------------|--|----------------------------------|--------------------------------------|--|--------------|--------------------------------------|--------------|--|--------------|---|------------------------------------|-----------------------------|--------------|-----------------------|--------------|--------------|--|-------------------------|--------------|----------------------------------|--------------|--------------------------------------|--------------|--------------|--------------|--------------|--|--------------|----------------------------------|--------------|--|--------------|
| Fold Change (ratio) | 0.246437105 | 0.090848213 | 0.091111614 | 0.000806583 | | 0.110855696 | | 0.121953593 | | 0.168279383 | | 0.187743879 | 0.218509986 | 0.221960648 | 0.22460642 | 0.228917694 | 0.237704503 | | 0.260055335 | 0.264360188 | 0.279338963 | 0.279947233 | 0.281207961 | 0.282705156 | 0.29408728 | 0.297990459 | 0.313634138 | | 0.323566748 | 0.32670528 | 0.36717276 | 0.415471413 | 0.407999585 |
| Fold Change | -4.06 | -11.01 | -10.98 | 10.77 | | -9.02 | • | -8.20 | | -5.94 | | -5.33 | 4.58 | 4.51 | -4.45 | -4.37 | 4.21 | | -3.85 | -3.78 | -3.58 | -3.57 | -3.56 | -3.54 | 3.40 | -3.36 | -3.19 | | -3.09 | -3.06 | -2.72 | -2.41 | -2.45 |
| Gene Name Gene Name melanoma inhihitory activity ras-related | GTP-binding protein 4b | actin, gamma 2, smooth muscle, enteric | calponin 1, basic, smooth muscle | myosin, heavy polypeptide 11, smooth | eukarvotic translation initiation factor 4 | gamma, 1 | myosin, heavy polypeptide 11, smooth | muscle | eukaryotic translation initiation factor 4 | gamma, 1 | laminin, alpha 3 (nicein (150kD), kalinin | (165kD), BM600 (150kD), epilegrin) | leiomodin 1 (smooth muscle) | | DKFZP586P1422 protein | | | laminin, beta 3 (nicein (125kD), kalinin | (140kD), BM600 (125kD)) | | myosin, light polypeptide kinase | | inositol polyphosphate-1-phosphatase | | | | | RNA-binding protein gene with multiple | splicing | myosin, light polypeptide kinase | | phosphatidic acid phosphatase type 2A BNA bindiog protein gene with multiple | splicing |
| er node 2761X <u>Genbank</u> | AA461365 | D00654 | D17408 | 200 | 2000 | AI380979 | | AF001548 | | AA156998 | | L34155 | X54162 | AA034289 | W72194 | D10667 | AA541622 | | U17760 | A1658662 | U48959 | AI768516 | L08488 | AI972873 | N95620 | AL043089 | C16443 | | D84110 | AA526844 | AB007972 | AF014402 | D84111 |
| 35 Genes from HCA cluster node 2761X SeqID Affy Genbank | 39271_at | 1197_at | 34203_at | 10 100 | 3/40/_s_all | 58774 at | ı | 767_at | | 63893_f_at | | 37909_at | 37765 at | 60532 at | 58409 at | 773 at | 64407 at | • | 36929 at | 50361 at | 32847 at | 62136_at | 41524 at | 65975 at | 57214 at | 65867 at | 50001 at | ı | 38048 at | 46276 at | 41137 at | 34797_at | 34162_at |
| 35 Genes SegID | 127 | 929 | 964 | 0 | 007 | 453 | | 283 | | 2 | | 1023 | 1223 | 17 | 1197 | 960 | 162 | | 1141 | 561 | 1151 | 658 | 1013 | 832 | 1099 | 869 | 926 | | 973 | 150 | 272 | 289 | 974 |
| Table 3; # | - | 7 | က | • | 4 | ß | | 9 | , | 7 | | œ | o | 10 | + | 12 | 13 | | 14 | 15 | 9 | 17 | 6 | 19 | 50 | 7 | 23 | | 23 | 24 | 52 | 56 | 27 |

| p-value | 2.789025E-04 | 5.432773E-06 | 1.694770E-05 | 3.251490E-06 | 1.715459E-03 | 4.653880E-06 | 4.038920E-05 | 3,551680E-04 |
|---------------------|-------------------|--------------|--------------|--------------|---------------------|-------------------|--------------|-----------------|
| Fold Change (ratio) | 0.389097343 | 0.381989673 | 0.473538178 | 0.47055836 | 0.458611832 | 0.348051542 | 0.367062886 | 0.406092748 |
| Fold Change | -2.57 | -2.62 | -2.11 | -2.13 | -2.18 | -2.87 | -2.72 | -2.46 |
| ene Name | | | | | | tein | | |
| Gene | | | | | CGI-43 protein | DKFZP564B0769 pro | | |
| Ø | | AI379892 | AI571525 | AI744109 | O | Δ | W87690 | W88427 |
| Genbank | AA843926 | | | | AI912571 C | W72919 D | | 54668_at W88427 |
| Affy Genbank G | 66529_at AA843926 | 49540_at | 49300_at | 65734 at | 46653_at AI912571 C | 49349 at W72919 D | | 54668_at |

| | Normal vs Stage III | 340.76+/-99.54 | N1=40, N2=10 | Fold Change: 2.01 | P-value: .00096 | 218.33+/-195.52 | 28.53+/-33.27 | N1=40, N2=10 | Fold Change: 5.14 | P-value: 0 | 252.16+/-278.17 | 15.58+/-46.81 | N1=39, N2=6 | Fold Change: 4.88 | P-value: 0 | | | × | | | 581.98+/-218.36 | 246.82+/-189 | N1=40, N2=10 | Fold Change: 2.76 | P-value: .00174 | 336.45+/-181.35 | 122.36+/-61.84 | N1=40, N2=10 | Fold Change: 2.82 | P-value: .0009 | 203.74+/-150.5 | -7.27+/-28.68 | N1=39, N2=6 | Fold Change: 6.91 | P-vaine: 0 | |
|---|---------------------|----------------|--------------|-------------------|-----------------|-----------------|----------------|---------------|-------------------|------------|-----------------|----------------|---------------|-------------------|-----------------|-----------------|----------------|---------------|-------------------|-----------------|-----------------|--------------|--------------|-------------------|-----------------|-----------------|----------------|--------------|-------------------|----------------|----------------|---------------|---------------|-------------------|-----------------|--|
| | Normal vs Stage II | | × | | | 218,33+/-195.52 | 49.83+/-71.53 | N1=40, N2=31 | Fold Change: 3.67 | P-value: 0 | 252.16+/-278.17 | 14.48+/-36.79 | N1=39, N2=31 | Fold Change: 4.51 | P-value: 0 | 221.4+/-284.63 | 86.7+/-207.7 | N1=40, N2=31 | Fold Change: 2.95 | P-value: .00006 | | | × | | | | | × | | | 203.74+/-150.5 | 41.75+/-90.59 | N1=39, N2=31 | Fold Change: 3.95 | P-value: 0 | |
| | Normal vs Stage I | | × | | | | | × | | | 252.16+/-278.17 | 47.94+/-61.86 | N1=39, N2=10 | Fold Change: 3.3 | P-value: .01164 | | | × | | | | | × | | | | | × | | | 203.74+/-150.5 | 56.05+/-43.37 | N1=39, N2=10 | Fold Change: 2.69 | P-value: .00466 | |
| OMA | Normal vs Malignant | | × | | | | | × | | | | | × | | | | | × | | | | | × | | | | | × | | | | | × | | | |
| Table 4: BREAST / INFILTRATING DUCT CARCINOMA | Normal vs All | | × | | | 218.33+/-195.52 | 75.87+/-104.51 | N1=40, N2=168 | Fold Change: 2.8 | P-value: 0 | 252.16+/-278.17 | 53.89+/-220.22 | N1=39, N2=168 | Fold Change: 3.64 | P-value: 0 | 221.16+/-280.96 | 57.63+/-138.17 | N1=40, N2=168 | Fold Change: 3.25 | P-value: 0 | | | × | | | | | × | | | 203.74+/-150.5 | 30.48+/-63.02 | N1=39, N2=168 | Fold Change: 4.19 | P-value: 0 | |
| ST / INFILTRA | Genbank | AA001250 | | | | AA017070 | | | | | AA027103 | | | | | AA029437 | | | | | AA029735 | | | | | AA031790 | | • | | | AA034289 | | | | | |
| Table 4: BREA | # Seq ID | 1 1 | | | | 7 2 | | | | | 3 11 | | | | | 4 12 | | | | | 5 13 | | | | | 6 15 | | | | | 7 17 | | | | | |

| | | | | | ** *********************************** | | |
|----|--------|----------|-------------------|---------------------|--|--------------------|---------------------|
| # | Seq ID | Genbank | Normal vs All | Normal vs Malignant | Normal vs Stage I | Normal vs Stage II | Normal vs Stage III |
| 8 | 18 | AA034418 | 370,77+/-175.09 | | | 367.92+/-176.43 | 367.92+/-176.43 |
| _ | | | 171.84+/-132.49 | | | 145.41+/-92.73 | 148.75+/-103.92 |
| | | | N1-40, N2=168 | × | × | N1=40, N2=31 | N1=40, N2=10 |
| | | | Fold Change: 2.41 | | | Fold Change: 2.76 | Fold Change: 2.55 |
| | | | P-value: 0 | | | P-value: 0 | P-value: .00125 |
| 6 | 20 | AA037766 | | | | 217.99+/-102.27 | |
| | | | | | | 98.41+/-70 | |
| | | | × | × | × | N1=39, N2=31 | × |
| | | | | | | Fold Change: 2.21 | |
| ٤ | 22 | AA044828 | 228.22+/-120.36 | 7000 | | 228.25+/-121.93 | 228.25+/-121.93 |
| 1 | ŀ | | 81.18+/-75.97 | | | 68.84+/-62.87 | 30.55+/-34.05 |
| | | | N1=40, N2=168 | × | × | NI=40, N2=31 | N1=40, N2=10 |
| | | | Fold Change: 2.87 | | | Fold Change: 3.18 | Fold Change: 5.8 |
| | | | P-value: 0 | | | P-value: 0 | P-value: 0 |
| Ξ | 23 | AA044830 | 384.63+/-189.59 | | | 387.92+/-190.91 | 387.92+/-190.91 |
| | | | 195.74+/-112.66 | | | 142.87+/-73.75 | 148.94+/-75.02 |
| | | | N1=40, N2=168 | × | × | N1=40, N2=31 | N1=40, N2=10 |
| | | | Fold Change: 2.09 | | | Fold Change: 2.81 | Fold Change: 2.65 |
| _ | | | P-value: 0 | | | P-value: 0 | P-value: .00182 |
| 12 | 24 | AA045145 | 261.46+/-178.02 | | | 262.21+/-180.28 | 262.21+/-180.28 |
| | | | 83.44+/-183.41 | | | 63.71+/-149.37 | 8.27+/-106.8 |
| | | | N1=40, N2=168 | × | × | N1=40, N2=31 | N1=40, N2=10 |
| _ | | | Fold Change: 3.35 | | | Fold Change: 4 | Fold Change: 5.08 |
| | | | P-value: 0 | | | P-value: 0 | P-value: .00018 |
| 2 | 25 | AA046457 | | | 254.96+/-154.86 | | 254.96+/-154.86 |
| | | | | | 91.58+/-30.86 | | 99.01+/-89.62 |
| | | | × | × | N1=40, N2=6 | × | N1=40, N2=10 |
| | | | | | Fold Change: 2.4 | | Fold Change: 2.75 |
| | | | | | P-value: .00032 | | P-value: .00204 |
| 14 | 27 | AA046853 | 200,27+/-196.52 | | 201.69+/-198.89 | 201.69+/-198.89 | 201.69+/-198.89 |
| | | | 385.33+/-229.71 | | 406.89+/-152.52 | 428.46+/-216.88 | 370.09+/-174.14 |
| | | | N1=40, N2=168 | × | N1=40, N2=6 | N1=40, N2=31 | N1=40, N2=10 |
| | | | Fold Change: 2.07 | | Fold Change: 2.42 | Fold Change: 2.44 | Fold Change: 2.06 |
| | | | P-value: 0 | | P-value: .0008 | P-value: 0 | P-value: .00306 |

| _ | | | | | | | | | _ | | | | _ | _ | | | _ | | _ | _ | | | _ | | | | | | | | _ | | _ |
|---------------------|----------------------------------|--------------|--------------------------------------|--------------|-----------------|--------------|-------------------|-----------------|----------------|----------------|--------------|-------------------|-----------------|-----------------|----------------|---------------|-------------------|-----------------|-----------------|-----------------|---------------|-------------------|-----------------|--------------|-----------------|---------------|-------------------|-----------------|-----------------|-----------------|--------------|-------------------|-----------------|
| Normal vs Stage III | 80.83+/-48.85 254.08+/-148.48 | N1=40, N2=10 | Fold Change: 3.07 P-value: .00013 | | | × | | | 282.9+/-254.52 | 62.96+/-63.59 | N1=40, N2=10 | Fold Change: 3.01 | P-value: .00335 | 383,25+/-127.97 | 164.81+/-71.68 | N1=40, N2=10 | Fold Change: 2.37 | P-value: .00006 | 450.12+/-413.33 | 4.41+/-98.3 | N1=40, N2=10 | Fold Change: 7.56 | P-value: 0 | 74.76+/-90.6 | 399.15+/-254.81 | N1=40, N2=10 | Fold Change: 5.85 | P-value: .00035 | 331.71+/-236.88 | 125.73+/-212.88 | N1=40, N2=10 | Fold Change: 4.1 | P-value: .00867 |
| Normal vs Stage II | | × | | 243+/-190.55 | 476.92+/-268.57 | N1=40, N2=31 | Fold Change: 2.22 | P-value: .00002 | 282.9+/-254.52 | 100.56+/-94.65 | N1=40, N2=31 | Fold Change: 2.24 | P-value: .00272 | 383.25+/-127.97 | 195.3+/-91.8 | N1-40, N2=31 | Fold Change: 2.03 | P-value: 0 | 450.12+/-413.33 | 78.85+/-105.76 | N1=40, N2=31 | Fold Change: 5.13 | P-value: 0 | 74.76+/-90.6 | 373,99+/-269.57 | N1=40, N2=31 | Fold Change: 5.56 | P-value: 0 | 331.71+/-236.88 | 176.14+/-256.71 | N1=40, N2=31 | Fold Change: 2.48 | P-value: .0012 |
| Normal vs Stage I | | × | | 243+/-190.55 | 413.4+/-148.7 | N1=40, N2=6 | Fold Change: 2.12 | P-value: .00235 | | | × | | | | | × | | | 450.12+/-413.33 | 84.82+/-108.93 | N1=40, N2=6 | Fold Change: 4.32 | P-value: .01621 | 74.76+/-90.6 | 328.32+/-224.02 | N1=40, N2=6 | Fold Change: 4.88 | P-value: .00774 | | | × | | |
| Normal vs Malignant | | × | | | | × | | | | | × | | | | | × | | | | | × | | | | | × | | | | | × | | |
| Normal vs All | | × | | | | × | | | | | × | | | 383.9+/-126.38 | 184.18+/-88.67 | N1=40, N2=168 | Fold Change: 2.2 | P-value: 0 | 444.15+/-409.74 | 108.09+/-137.58 | N1=40, N2=168 | Fold Change: 3.84 | P-value: 0 | 73.64+/-89.7 | 421.17+/-253.37 | N1=40, N2=168 | Fold Change: 6.85 | P-value: 0 | | | × | | |
| Genbank | AA056180 | | | AA056755 | | | | | AA058578 | | | | | AA059396 | | | | | AA059401 | | | | | AA059458 | | | | | AA075632 | | | | |
| Seq ID | 78 | | | 53 | | | | | 30 | | | | | 31 | | | | | 32 | | | | | 33 | | | | | 34 | | | | |
| * | 15 | | | 16 | | | | | 17 | | | | | 18 | | | | | 19 | | | | | 20 | | | | | 21 | | | | |

| # | Seq ID | Genbank | Normal vs All | Normal vs Malignant | Normal vs Stage I | Normal vs Stage II | Normal vs Stage III |
|----|--------|----------|-------------------|---------------------|-------------------|--------------------|---------------------|
| 77 | 35 | AA079839 | | | | 399.21+/-217.44 | 399.21+/-217.44 |
| | | | | | | 178.24+/-173.52 | 173.1+/-209.18 |
| | | | × | × | × | N1=39, N2=31 | N1=39, N2=6 |
| | | | | | | Fold Change: 2.2 | Fold Change: 2.61 |
| | | | | | | P-value: .00001 | P-value: .00239 |
| 23 | 37 | AA082546 | 272.4+/-223.42 | | | 275.39+/-225.53 | 275.39+/-225.53 |
| | | | 105.34+/-63.79 | | | 93.91+/-47.44 | 74.83+/-36.32 |
| | | | N1=40, N2=168 | × | × | N1=40, N2=31 | N1=40, N2=10 |
| | | | Fold Change: 2.35 | | | Fold Change: 2.55 | Fold Change: 3.23 |
| | | | P-value: 0 | | | P-value: 0 | P-value: .00005 |
| 24 | 38 | AA101125 | | | | 670.66+/-337.66 | 670.66+/-337.66. |
| | | | | | | 325+/-154.53 | 292.03+/-126.99 |
| | | | × | × | × | N1=40, N2=31 | N1=40, N2=10 |
| | | | | | | Fold Change: 2.01 | Fold Change: 2.27 |
| | | | | | | P-value: 0 | P-value: .00091 |
| 23 | 41 | AA126704 | 314.32+/-135.99 | | | 312.64+/-137.34 | 312,64+/-137,34 |
| | | | 107,92+/-81.49 | | | 77.02+/-71.72 | 78.95+/-75.5 |
| | | | N1=40, N2=168 | × | × | N1-40, N2-31 | N1=40, N2=10 |
| | | | Fold Change: 3.17 | | | Fold Change: 4.14 | Fold Change: 3.75 |
| | | | P-value: 0 | | | P-value: 0 | P-value: .00032 |
| 56 | 42 | AA127718 | 238.22+/-357.2 | | | 240.21+/-361.64 | |
| | | | 82.46+/-76.05 | | | 87.69+/-80.58 | |
| | | | N1=40, N2=168 | × | × | N1=40, N2=31 | × |
| | | | Fold Change: 2.53 | | | Fold Change: 2.39 | |
| | | | P-value: 0 | | | F-value: .00006 | |
| 27 | 43 | AA127727 | 214.24+/-122.15 | | 212.97+/-123.48 | 212.97+/-123.48 | 212.97+/-123.48 |
| | | | 81.63+/-51.81 | | 89.92+/-52.2 | 66.11+/-40.88 | 51.85+/-29.44 |
| | | | N1=40, N2=168 | × | N1=40, N2=6 | N1=40, N2=31 | N1=40, N2=10 |
| | | | Fold Change: 2.72 | | Fold Change: 2.29 | Fold Change: 3.29 | Fold Change: 3.99 |
| | | | P-value: 0 | | P-value: .00795 | P-value: 0 | P-value: .00001 |
| 28 | 47 | AA131456 | 642,53+/-392,68 | | | 650.36+/-394.64 | |
| | | | 325.78+/-211.65 | | | 340.69+/-188.84 | |
| | | | N1=40, N2=168 | × | × | N1=40, N2=31 | × |
| | | | Fold Change: 2.13 | | | Fold Change: 2.01 | |
| | | | P-value: 0 | | | P-value: 0 | |

| * | Seq ID | Genbank | Normal vs All | Normal vs Malignant | Normal vs Stage I | Normal vs Stage II | Normal vs Stage III |
|----|--------|------------|---------------------------------|--|-------------------|---|---------------------------------|
| 29 | 48 | AA131632 | | | | 376.43+/-162.83 185.82+/-88.41 | |
| | | | × | × | × | N1=40, N2=31 Fold Change: 2.11 P-value: 0 | × |
| 98 | 20 | AA131894 | | | | | 125.64+/-46.19 |
| | | | × | × | × | × | N1-40, N2-10 |
| | | | | | | | Fold Change: 2.19 |
| 12 | ū | A A 122748 | AD1 80±7-133 1A | | | | 400 01+/-134 73 |
| 5 | 5 | 0.00000 | 193.81+/-109.33 | | | | 182.12+/-50.15 |
| | | | N1=40, N2=168 | × | × | × | N1=40, N2=10 |
| | | | Fold Change: 2.26 P-value: 0 | | | | Fold Change: 2.15 P-value: 0 |
| 32 | 55 | AA137038 | | | | 122.53+/-86.86 | |
| | | | | | | 79.507-/-10.007 | 1 |
| | | | × | × | × | N1=40, N2=31 | × |
| | | | | | | P-value: .00008 | |
| 33 | 57 | AA142913 | 302,75+/-219,97 | | 302.34+/-222.83 | 302.34+/-222.83 | 302,34+/-222.83 |
| | | | 88.13+/-60.53 | | 104.62+/-49.26 | 84.31+/-62.54 | 83.75+/-83.82 |
| | | | N1=40, N2=168 | × | N1=40, N2=6 | N1=40, N2=31 | N1=40, N2=10 |
| | | | Fold Change: 3.49 | | Fold Change: 2.6 | Fold Change: 3.74 | Fold Change: 4.01 |
| | | | P-value: 0 | | P-value: .00174 | P-value: 0 | P-value: .00014 |
| 34 | 65 | AA143491 | 491.76+/-382.75 | | | | 485.72+/-385.82 |
| | | | 278.14+/-268.9 | | | | 206.94+/-269.35 |
| | | | N1=40, N2=168 | × | × | × | NI=40, N2=10 |
| | | | Fold Change: 2.1 | | | | Fold Change: 3.02 |
| | | | P-value: .00012 | | | | P-value: .02186 |
| 35 | 62 | AA147751 | 480.16+/-205.12 | | | 478.2+/-207.42 | |
| | | | 247.19+/-214.23 | | | 284.09+/-237.65 | |
| | | | N1=40, N2=168 | × | × | N1=40, N2=31 | × |
| | | | Fold Change: 2.38 | | | Fold Change: 2.21 | |
| | | | P-value: 0 | and the second s | | P-value: .00015 | |

| | | | | | 1 | | |
|------|--------|----------|--|---------------------|--|---|---|
| * | Sed ID | Genbank | Normal vs All | Normal vs Malignant | Normal vs Stage I | Normal vs Stage II | Normal vs Stage III |
| 98 | 63 | AA147884 | 46.57+/-54.48 235.82+/-200.06 N1=40, N2=168 Fold Change: 4.33 P-value: 0 | × | × | × | × |
| 37 | 2 | AA149312 | 377.26+/-139.17 193.77+/-99.25 N1=40, N2=168 Fold Change: 2.04 .P-value: 0 | × | × | 374+/-139,43 170,47+/-78.83 N1=40, N2=31 Fold Change: 2.29 P-value: 0 | 374+/-139.43 140.83+/-37.98 N1=40, N2=10 Fold Change: 2.57 P-value: 0 |
| ee . | 89 | AA150501 | 213.29+/-103.88 89.09+/-51.7 N1=40, N2=168 Fold Change: 2.45 P-value: 0 | × | 215.8+/-104 97.8+/-28.03 N1=40, N2=6 Fold Change: 2.01 P-value: .00034 | 215.8+/-104 74.57+/-38.51 N1=40, N2=31 Fold Change: 2.85 P-value: 0 | 215.8+/-104 41.35+/-30.37 N1=40, N2=10 Fold Change: 4.75 P-value: 0 |
| £ | 99 | AA151346 | × | × | × | 180.8+/-85.07 392.04+/-178.92 N1=39, N2=31 Fold Change: 2.13 P-value: 0 | × |
| 40 | 89 | AA155914 | 400.19+/405.95 143.7+/-119.6 N1=40, N2=168 Fold Change: 2.56 P-value: 0 | × | × | 407.07+/408.89 135.07+/-104.23 N1=40, N2=31 Fold Change: 2.69 P-value: .00009 | × |
| 4 | જ | AA155952 | × | × | × | 343.72+/-239.12 136.91+/-62.46 N1=40, N2=31 Fold Change: 2.23 P-value: 0 | 343.72+1/239.12 111.1+1/-55.68 N1=40, N2=10 Fold Change: 2.77 P-value: .00005 |
| 24 | IT. | AA158731 | 288.7+/-238.19 93.33+/-144.72 N1=40, N2=168 Fold Change: 3.48 P-value: 0 | x | × | 287.72+/-241.22 95.77+/-117.6 N1=40, N2=31 Fold Change: 3.08 P-value: .00001 | 287.72+/-241.22 183.13+/-413.53 N1=40, N2=10 Fold Change: 3.9 P-value: .01612 |
| | | | | | | | |

| # | Seq ID | Genbank | Normal vs All | Normal vs Malignant | Normal vs Stage I | Normal vs Stage II | Normal vs Stage III |
|----|--------|----------|-------------------|---------------------|-------------------|--------------------|---------------------|
| 64 | 73 | AA165701 | 176,56+/-116,87 | | 176.56+/-116.87 | 176.56+/-116.87 | 176.56+/-116.87 |
| | ! | | 335.74+/-147.31 | | 322.48+/-173.1 | 299.01+/-131.22 | 309.98+/-142.67 |
| | | | N1-40, N2=168 | × | N1=40, N2=6 | N1=40, N2=31 | N1=40, N2=10 |
| | | | Fold Change: 2.21 | | Fold Change: 2.14 | Fold Change: 2.01 | Fold Change: 2.05 |
| | | | P-value: 0 | | P-value: .0095 | P-value: .00002 | P-value: .00163 |
| 4 | 74 | AA166620 | 227.06+/-165.92 | | 227.06+/-165.92 | 227.06+/-165.92 | 227.06+/-165.92 |
| | | | 100.03+/-194.31 | | 88.27+/-25.59 | 85.25+/-119.41 | 38.81+/-22.68 |
| | | | N1=39, N2=168 | × | N1=39, N2=10 | N1=39, N2=31 | N1=39, N2=6 |
| | | | Fold Change: 2.58 | | Fold Change: 2.06 | Fold Change: 2.98 | Fold Change: 4.89 |
| | | | P-value: 0 | | P-value: .00143 | P-value: 0 | P-value: 0 |
| 45 | 27 | AA173572 | 368.06+/-171.39 | | | 368.73+/-173.58 | 368.73+/-173.58 |
| ! | ! | | 132.7+/-121.15 | | | 105.87+/-79.3 | 162.1+/-222.38 |
| | | | N1=40, N2=168 | × | × | N1=40, N2=31 | N1=40, N2=10 |
| | | | Fold Change: 3.31 | | | Fold Change: 3.94 | Fold Change: 3.72 |
| | | | P-value: 0 | | | P-value: 0 | P-value: .0091 |
| 46 | 11 | AA188763 | 109.78+/-67.46 | | | | 109.94+/-68.33 |
| : | : | | 223,51+/-105,81 | | | | 210.03+/-82.61 |
| | | | N1=40, N2=168 | × | × | × | N1=40, N2=10 |
| | | | Fold Change: 2.13 | | | | Fold Change: 2.06 |
| | | | P-value: 0 | | | | P-value: .00159 |
| 7 | 78 | AA193340 | | | | 338.82+/-318.56 | 338.82+/-318.56 |
| : | | | | | | 135.54+/-138.52 | 65.09+/-44.54 |
| | | | × | × | × | N1=40, N2=31 | N1=40, N2=10 |
| | | | | | | Fold Change: 2.47 | Fold Change: 4.45 |
| | | | | | | P-value: .00001 | P-value: .00006 |
| 84 | 85 | AA211369 | | | | | 495.8+/-374.32 |
| | | | | | | | 166.56+/-142.09 |
| | | | × | × | × | × | N1=40, N2=10 |
| | | | | | | | Fold Change: 2.75 |
| | | | | | | | P-value: .00161 |
| 6 | 88 | AA227778 | 250.35+/-164.31 | | | 254,32+/-164.5 | 254.32+/-164.5 |
| | | | 103.23+/-228.68 | | | 157,31+/-463.97 | 92.7+/-116.55 |
| | | | N1=40, N2=168 | × | × | N1=40, N2=31 | N1=40, N2=10 |
| | | | Fold Change: 3.07 | | | Fold Change: 3.09 | Fold Change: 2.93 |
| | | | P-value: 0 | | | P-value: .00002 | P-value: .00666 |

| # Seq ID Geabank Normal v Millgmant Normal v Stage II Normal v Stage III 9 9 5 | | | | | | | , | |
|--|-----|--------|----------|-------------------|---------------------|-------------------|---|---------------------|
| 95 AA296461 X X X X TABABABABABABABABABABABABABABABABABABAB | # | Seq ID | Genbank | Normal vs All | Normal vs Malignant | Normal vs Stage I | Normal vs Stage II | Normal vs Stage III |
| National Changes 205 | 20 | 35 | AA295451 | | | | 428.38+/-345.54 191.85+/-157.04 | |
| 56 AA310786 X | | | | × | × | × | N1=40, N2=31 Fold Change: 2.05 | × |
| 96 AA310786 24.707+10.49 | | | | | | | P-value: .0027 | |
| National Changes 2.19 No.2547-1514 | 12 | 96 | AA310786 | | | | 242.07+/-109.49 | 242.07+/-109.49 |
| National Color National Color | | | | | | | 509.52+/-191.41 | 578.59+/-173.41 |
| Fording 2.19 | | | | × | × | × | N1=40, N2=31 | N1=40, N2=10 |
| 97 AA312965 X X X 570,534-105.99 55404-1183.8 56404-1183.8 56404-1183.8 56404-1183.8 56404-1183.8 56404-1183.8 56404-1183.8 56404-1183.8 56404-1183.8 56404-1183.8 56404-1183.8 56404-1183.8 56404-1183.8 56404-1183.8 56404-1183.8 56404-1183.9 56404-1183.8 56404-1183. | | | | | | | Fold Change: 2.19 | Fold Change: 2.55 |
| 97 AA312065 X X X X X X X X X X X X X X X X X X X | | | | | | | P-value: 0 | P-value: 0 |
| National Control Con | 22 | 26 | AA312905 | | | | 270.53+/-105.99 | |
| National Color National Color | | | | | | | 554.04+/-183.58 | |
| Post Change: 2.06 | | | | × | × | × | N1=40, N2=31 | × |
| 99 AA369887 X X X X X X X X X X X X X X X X X X | | | | | | | Fold Change: 2.06 | |
| 102 AA400000 X | lg | g | 78893FAA | | | | | 326.24+/-259.48 |
| 102 | 2 | ; | | | | | | 88 98+/-90 46 |
| 102 AA400080 X X X X X X X X X | | | | Þ | Þ | Þ | Þ | N1=40 N2=10 |
| 102 AA400680 X X X X X X X X X | | | | < | < | < | < | Told Change, 4 05 |
| 10.2 AA410680 X | | | | | | | | P-value: .00073 |
| Name | 4 | 102 | AA400080 | | | | 89.66+/-112.79 | |
| Name | | | | | | | 246.79+/-233.75 | |
| 104 AA47'813 846.09H-411.44 Paches, 221 Paches, 221 Paches, 221 Paches, 221 Paches, 222 | | | | × | × | × | N1=40, N2=31 | × |
| 104 AA417813 846.09+ -411.44 Propher, 0.045 STINGE-LEGALS STINGE-LEGALS Standard Propher, 0.045 N1=64, Par-18, P | | | | | | | Fold Change: 2.21 | |
| 104 AA417813 846.09+411.44 849.01+4.416.4 104 AA417813 87.08+6.24.59 X N1=40, N2=51.64 105 AA418636 N1=40, N2=16 N1=40, N2=10 106 AA418636 N1=40, N2=10 N1=40, N2=10 107 AA418636 N1=40, N2=10 N1=20, N2=10 108 AA418636 N1=40, N2=10 N1=20, N2=10 109 N2=10 N1=20, N2=10 109 N2=10 N1=20, N2=10 109 N2=10 N2=10 N1=20, N2=10 109 N2=10 N1=20, N2=10 109 N2=10 N2=10 N2=10 N2=10 N2=10 N2=10 109 N2=10 N2=10 N2=10 N2=10 N2=10 109 N2=10 N2=10 N2=10 N2=10 N2=10 N2=10 109 N2=10 N2= | | | | | | i | P-value: .0043 | |
| NI=di, NN=168 | lik | 104 | AA417813 | 846.09+/-411.44 | | | 849.01+/-416.4 | |
| N N N N N N N N N N | | | | 570.86+/-624.96 | | | 499.89+/-596.68 | |
| Fold Change: 2.04 Fold Change: 2.44 Fold Change: 2.44 Foreibre: (20007) Fore | | | | N1=40, N2=168 | × | × | NI=40, N2=31 | × |
| P-valine: 0 P-valine: 0 P-valine: 0.0007 | | | | Fold Change: 2.01 | | | Fold Change: 2.44 | |
| 106 | | | | P-value: 0 | | | P-value: .00007 | |
| X N1=39, N2=10 N1=39, N2=10 N1=39, N2=31 Fold Change: 2.43 Fold Change: 3.75 P-value: .0032 P-value: 0 | 26 | 106 | AA418636 | 244.96+/-112.2 | | 244.96+/-112.2 | 244.96+/-112.2 | 244.96+/-112.2 |
| X N1=59, N2=10 N1=59, N2=51 Fold Change: 243 Fold Change: 3.75 F.value: .0032 F.value: 0 | | | | 85.16+/-53.2 | | 107.39+/-80.98 | 62.82+/-36.83 | 57.15+/-13.69 |
| Fold Change: 2.43 Fold Change: 3.75 P-value: .02032 P-value: 0 | | | | N1=39, N2=168 | × | N1=39, N2=10 | N1=39, N2=31 | N1=39, N2=6 |
| P-value: .02032 P-value: 0 | | | | Fold Change: 2.92 | | Fold Change: 2.43 | Fold Change: 3.75 | Fold Change: 3.86 |
| | | | | P-value: 0 | | P-value: .02032 | P-value: 0 | P-value: 0 |

| State Colonial C | 1 | 8 | Contract. | M* | Manager Manager | T | M | M |
|---|----|--------|-----------|-------------------|--------------------|-------------------|--------------------|---------------------|
| 107 AA418995 | # | Sed ID | Genbank | Normal vs All | Normal vs Mangnant | Normal Vs Stage I | Normal VS Stage II | Normal vs Stage III |
| 109 AAA19263 X | 57 | 107 | AA418995 | | | | | 276.52+/-180.68 |
| 109 AAA19263 | | | | | | | | 89.94+/-34.01 |
| 109 AAA19263 X | | | | × | × | × | × | N1=40, N2=10 |
| 109 AAA19263 | | | | | | | | Fold Change: 2.64 |
| 199 AAA19263 X | | | | | | | | P-value: 0 |
| 112 AAA43460 72764-123.56 73.834+124.68 73.834+124.68 73.834+124.68 73.834+124.68 73.834+124.68 73.834+124.68 73.834+124.68 73.834+124.68 73.834+124.68 73.834+124.68 73.834+124.68 73.834+124.68 73.834+124.68 73.834+124.68 73.834+124.68 73.834+124.23 73.834+123.24 73.834+123.2 | 85 | 109 | AA419263 | | | | | 547.83+/-148.39 |
| 112 AA424160 72.764-123.56 73.834-124.68 73.834-124.68 73.834-124.68 73.834-124.68 73.834-124.68 73.834-124.68 73.834-124.68 73.834-124.68 73.834-124.68 73.834-124.68 73.834-124.68 73.834-124.68 73.834-124.68 73.834-124.68 73.834-124.68 73.834-124.69 73.834-124. | | | | | | | | 268.37+/-114.78 |
| 11.2 AA434160 72.764-123.26 73.834-124.68 73.834-124.68 73.834-124.68 73.834-124.68 73.834-124.68 73.834-124.68 73.834-124.68 73.834-124.68 73.834-124.68 73.834-124.69 73.834-124.22 73.834-124.23 73.834-124 | | | | × | × | × | × | N1=40, N2=10 |
| 112 AA424160 72.764-123.26 73.824-124.68 341.94-203.22 342.64-37.34 341.94-203.22 342.64-37.34 341.94-203.22 342.64-37.34 341.94-203.22 342.64-37.34 341.94 32.94.64.20 32.84-144.20 32.84-144.20 32.84-144.20 32.84-144.20 32.84-144.20 32.84-144.20 32.84-144.20 32.84-144.20 32.84-144.20 32.84-144.20 32.84-144.20 32.84-144.20 32.84-144.20 32.84-144.20 32.84-144.20 32.84-144.20 32.84-14.20 | | | | | | | | Fold Change: 2.12 |
| 112 | | | | | | | | P-value: .00021 |
| National Change 2.08 National Change 2.09 National Change 2.04 National Change 2.47 National Change 2.48 National Change 2.48 | 59 | 112 | AA424160 | 72.76+/-123.26 | | | 73.82+/-124.68 | 73.82+/-124.68 |
| N1-40, N2-168 | | | | 241.99+/-260.32 | | | 344.26+/-347.34 | 405.04+/-245.91 |
| Fold Changes 3.39 Fold Changes 5.08 | | | | N1=40, N2=168 | × | × | N1=40, N2=31 | N1=40, N2=10 |
| 115 | | | | Fold Change: 3.39 | | | Fold Change: 5.08 | Fold Change: 7.49 |
| 115 | | | | P-value: 0 | | | P-value: 0 | P-value: 0 |
| 124.83+1.150.06 | 8 | 115 | AA429308 | 242.38+/-144.29 | | | 237.83+/-143.23 | 237.83+/-143.23 |
| M1-40, N2-168 | | | | 124.83+/-150.06 | | | 88,15+/-43.75 | 65.39+/-41.56 |
| Fold Change 2.04 Fold Change 2.47 | | | | N1=40, N2=168 | × | × | N1=40, N2=31 | N1=40, N2=10 |
| 116 | | | | Fold Change: 2,04 | | | Fold Change: 2.47 | Fold Change: 3.5 |
| 116 | | | ï | P-value: 0 | | | P-value: 0 | P-value: ,00009 |
| 117 AAA30314 265.86F+187.91 259.57F+186.05 118 AAA430185 N1-40, N2-168 X X X X X X X X X | 12 | 116 | AA430300 | | | | | 511.45+/-201.77 |
| 117 AAA30134 265.86F-187.91 259.57F-186.05 259. | | | | | | | | 242.83+/-223.58 |
| 117 AAA30314 265.86F/187.91 259.57F/186.05 257F/186.05 257F/ | | | | × | × | × | × | N1=40, N2=10 |
| 117 AAA30314 265.864-187.91 259.574-101.82 259. | | | | | | | | Fold Change: 2.55 |
| 117 | | | | | | | | P-value: .00275 |
| Mark | 62 | 117 | AA430314 | 265.86+/-187.91 | | | 259.57+/-186.05 | 259.57+/-186.05 |
| N1-40, N2-168 | | | | 65.06+/-109.36 | | | 59.77+/-101.82 | -4.03+/-59 |
| Fold Changer 3.88 Fold Changer 4.16 P-value: 0 | | | | N1=40, N2=168 | × | × | NI=40, N2=31 | N1=40, N2=10 |
| 118 | | | | Fold Change: 3.88 | | | Fold Change: 4.16 | Fold Change: 6.66 |
| 118 AAA30185 233,344-335.89 710,154-40,136 710,154-40,136 710,154-40,136 710,154-40,136 710,154-40,136 710,1440,136-238 710,1440,1460,1460,1460,1460,1460,1460,14 | | | | P-value: 0 | | | P-value: 0 | P-value: 0 |
| X X X X X X X X X X X X X X X X X X X | 8 | 118 | AA436185 | | | | 253.53+/-335.89 | |
| A A N.—3.1 Fold Change: 2.38 P-value: 00089 | | | | Þ | Þ | Þ | 70.15+/-61.86 | Þ |
| FOR CARDING AND A CONTROL AND | | | | 4 | 4 | 4 | Todd Change 220 | < |
| | | | | | | | P-resine: 00080 | |

| # | Seq 10 | Genbank | Normal vs All | Normal vs Malignant | Normal vs Stage I | Normal vs Stage II | Normal vs Stage III |
|----|--------|------------|-------------------|---------------------|-------------------|--------------------|---------------------|
| 64 | 119 | AA436250 | | | | 213.31+/-226.04 | |
| | l | | | | | 84,53+/-80,67 | |
| | | | × | × | × | N1=40, N2=31 | × |
| | | | | | | Fold Change: 2.03 | |
| | | | | | | P-value: .00323 | |
| ŝ | 120 | AA447015 | 227.33+/-171.55 | | | 226.67+/-173.74 | |
| | | | 82.62+/-74.49 | | | 66.02+/-51.23 | |
| | | | N1=40, N2=168 | × | × | N1=40, N2=31 | × |
| | | | Fold Change: 2.51 | | | Fold Change: 2.87 | |
| | | | P-value: 0 | | | P-value: 0 | |
| 99 | 122 | AA450090 | 285.47+/-226.15 | | | 285.47+/-226.15 | |
| | | | 133,85+/-131,49 | | | 125.82+/-152.5 | |
| | | | N1=40, N2=168 | × | × | N1=40, N2=31 | × |
| | | | Fold Change: 2.57 | | | Fold Change: 2.88 | |
| | | | P-value: 0 | | | P-value: 0 | |
| 29 | 124 | A A 452295 | 227.07+/-122.52 | | 220,36+/-116.43 | 220.36+/-116.43 | 220.36+/-116.43 |
| | | | 44.21+/-53.65 | | 97.66+/-165.2 | 39.86+/-35.12 | 45.22+/-26.99 |
| | | | N1=40, N2=168 | × | N1=40, N2=6 | N1=40, N2=31 | N1=40, N2=10 |
| | | | Fold Change: 5.1 | | Fold Change: 3.73 | Fold Change: 5.14 | Fold Change: 4.25 |
| | | | P-value: 0 | | P-value: .03669 | P-value: 0 | P-value: 0 |
| 8 | 125 | AA455877 | | | | 295.74+/-130.76 | 295.74+/-130.76 |
| } | | | | | | 131.5+/-76.16 | 102.29+/-56 |
| | | | × | × | × | N1=40, N2=31 | N1=40, N2=10 |
| | | | | | | Fold Change: 2,4 | Fold Change: 3.08 |
| | | | | | | P-value: 0 | P-value: .0007 |
| 9 | 126 | AA456099 | | | | | 375.25+/-161.56 |
| | İ | | | | | | 163.63+/-115.97 |
| | | | × | × | × | × | N1=40, N2=10 |
| | | | | | | | Fold Change: 2.5 |
| | | | | | | | P-value: .00094 |
| 6 | 128 | AA464792 | | | | | 244.42+/-163.62 |
| | | | | | | | 77.5+/-57.57 |
| | | | × | X | × | × | N1=40, N2=10 |
| | | | | | | | Fold Change: 2.92 |
| | | | | | | | P-value: .00105 |

| # | Sed ID | Genbank | Normal vs All | Normal vs Malignant | Normal vs Stage I | Normal vs Stage II | Normal vs Stage III |
|--------|--------|----------|--|---------------------|-------------------|--|--|
| Ę | 129 | AA479033 | 104.14/-260.94 492.384/-952.15 N1=40, N2=168 Fold Change: 2.62 P-value: .00001 | × | × | × | × |
| t 1 | 131 | AA480075 | 331,51+/-157,28 155,12+/-117,94 N1=40, N2=168 Fold Change: 2.34 P-value: 0 | × | × | 331,54/-159.34 119.29+/-99.73 N1=40, N2=31 Fold Change: 3.04 P-value: 0 | 331.5+/-159.34 97.24+/-36.96 N1=40, N2=10 Fold Change: 3.11 P-value: 0 |
| 57 | 133 | AA486366 | 237.55+/-149.6 115.8+/-122.37 N1=39, N2=168 Fold Change: 2.09 P-value: 0 | × | X | 237.554/-149.6 102.264/-105.5 N1=39, N2=31 Fold Change: 2.18 P-value: .0001 | 237,554-149.6 20,974/-35.35 N1=39, N2=6 Fold Change: 4.64 P-value: 0 |
| 4. | 135 | AA488889 | 298.27+/-192.46 99.12+/-55.38 N1=40, N2=168 Fold Change: 2.64 P-value: 0 | × | × | 298.86+/-194.94 90.94+/-37.43 N1=40, N2=31 Fold Change: 2.8 P-value: 0 | 298.86+/-194.94 118.43+/-135.55 N1=40, N2=10 Fold Change: 2.62 P-value: .00133 |
| 85 | 136 | AA496142 | × | × | × | 223+/-72.35 113.96+/-51.66 N1=39, N2=31 Fold Change: 2.03 P-yalue: 0 | × |
| 92 | 137 | AA501987 | 202.154/-87.72 123.464/-65.5 N1=39, N2=168 Fold Change: 1.68 P-value: 0 | × | × | × | × |
| 12 | 138 | AA502943 | 438.99+/-109.54 262.83+/-186.3 N1=40, N2=168 Fold Change: 2.11 P-value: 0 | × | × | 439.24+/-110.96 250.93+/-145.84 N1=40, N2=31 Fold Change: 2.17 P-value: .00002 | × |

| * | Seq ID | Genbank | Normal vs All | Normal vs Malignant | Normal vs Stage I | Normal vs Stage II | Normal vs Stage III |
|----|--------|------------|-------------------|---------------------|-------------------|--------------------|---------------------|
| 28 | 139 | AA504253 | | | | | 394.39+/-105.44 |
| | | | × | × | × | × | N1=40, N2=10 |
| | | | | | | | Fold Change: 2.81 |
| | | | | | | | P-value: .00011 |
| 79 | 140 | AA508196 | 475.57+/-315.6 | | | 475.57+/-315.6 | 475.57+/-315.6 |
| | | | 223.41+/-148.84 | | | 207.81+/-174.35 | 184.11+/-120.65 |
| | | | N1=40, N2=168 | × | × | N1=40, N2=31 | N1=40, N2=10 |
| _ | | | Fold Change: 2.02 | | | Fold Change: 2.39 | Fold Change: 2.51 |
| | | | P-value: 0 | | | P-value: .00002 | P-value: .00343 |
| 8 | 141 | AA513002 | 156.4+/-98.44 | | | 156.4+/-98.44 | |
| | | | 285.99+/-134.83 | | | 282.09+/-129.91 | |
| | | | N1=39, N2=168 | × | × | N1=39, N2=31 | × |
| | | | Fold Change: 2.05 | | | Fold Change: 2.03 | |
| 6 | 146 | A A 524095 | T THIRD: O | | | | 1037.85+/-287.66 |
| 3 | | | | | | | 517.53+/-221.07 |
| | | | × | × | × | × | N1=40, N2=10 |
| | | | 1 | | | | Fold Change: 2.08 |
| | | | | | | | P-value: .00018 |
| 8 | 147 | A A 524250 | 337.9+/-133.73 | | | 340.18+/-134.68 | 340.18+/-134.68 |
| ! | | | 180.07+/-116.79 | | | 149.6+/-62.5 | 98.23+/-47.07 |
| | | | N1=40, N2=168 | × | × | N1=40, N2=31 | N1=40, N2=10 |
| | | | Fold Change: 2.02 | | | Fold Change: 2.28 | Fold Change: 3.58 |
| | | | P-value: 0 | | | P-value: 0 | P-value: .00001 |
| 83 | 148 | AA524536 | 304.34+/-210.76 | | 298.68+/-210.41 | 298.68+/-210.41 | |
| | | | 116.11+/-126.7 | | 87.05+/-113.19 | 113.27+/-115.65 | |
| | | | N1=40, N2=168 | × | N1-40, N2-6 | N1=40, N2=31 | × |
| | | | Fold Change: 2.64 | | Fold Change: 3 | Fold Change: 2.42 | |
| | | | P-value: .00001 | | P-value: .03651 | P-value: .00101 | |
| 8 | 151 | AA526961 | 415,49+/-234,41 | | | 417.14+/-237.24 | 417.14+/-237.24 |
| | | | 178.69+/-118.79 | | | 163.41+/-149.82 | 119.79+/-62.48 |
| | | | N1=40, N2=168 | × | × | N1=40, N2=31 | N1=40, N2=10 |
| | | | Fold Change: 2.31 | | | Fold Change: 2.62 | Fold Change: 3.37 |
| | | | P-value: 0 | | | P-value: 0 | P-value: .0000/ |

| 3 | Genbank | Normal vs All | Normal vs Malignant | Normal vs Stage I | Normal vs Stage II | Normal vs Stage III |
|----------|--------------|---|---------------------|-------------------|--|---|
| AA527180 | | X | × | × | × | 336.44+/-180.05 159.84+/-85.84 N1=40, N2=10 Fold Change: 2.03 P-value: .00507 |
| AA527298 | | 121.15+/-82.59 287.83+/-242.33 N1=39, N2=168 Fold Change: 1.92 P-value: 0 | × | × | × | X |
| AA533272 | | × | × | × | 345.7+/-116.17 171.66+/-72 N1=40, N2=31 Fold Change: 2.1 P-value: 0 | × |
| AA534724 | * | × | × | × | 143.93+/-57.72 325.98+/-117.74 N1=40, N2=31 Fold Change: 2.31 P-value: 0 | 143.93+/-57.72 353.03+/-109.1 N1=40, N2=10 Fold Change: 2.56 P-value: 0 |
| AA535218 | _∞ | 323.08+/-135.81 155.95+/-97.92 N1=40, N2=168 Fold Change: 2.27 P-value: 0 | × | × | 322.09+/-137.43 178.24+/-126.11 N1=40, N2=31 Fold Change: 2.06 P-value: .00002 | × |
| AA543076 | يو | × | × | × | 109.8+/-95.15 365.31+/-441.38 N1=40, N2=31 Fold Change: 2.58 P-value: .00003 | × |
| AA555312 | 7 | × | × | × | × | 215.48+/-192.29 40.38+/-31.93 N1=40, N2=10 Fold Change: 3.68 P-value: .00006 |

| * | Seq ID | Genbank | Normal vs All | Normal vs Malignant | Normal vs Stage I | Normal vs Stage II | Normal vs Stage III |
|----|--------|----------|-------------------|---------------------|-------------------|--------------------|---------------------|
| 6 | 169 | AA568397 | 243.4+/-227.66 | | 243.4+/-227.66 | 243.4+/-227.66 | 243.4+/-227.66 |
| | | | 92.45+/-68.55 | | 86.44+/-37.31 | 74.95+/-70.23 | 52.06+/-33.54 |
| | | | N1=40, N2=168 | × | N1=40, N2=6 | N1=40, N2=31 | N1=40, N2=10 |
| | | | Fold Change: 2.27 | | Fold Change: 2.12 | Fold Change: 2.77 | Fold Change: 3.85 |
| | | | P-value: 0 | | P-value: .00208 | P-value: 0 | P-value: .00005 |
| 93 | 170 | AA570519 | 200.18+/-172.78 | | | | 200.18+/-172.78 |
| _ | | | 77.04+/-65.26 | | | | 48.77+/-33 |
| | | | N1=40, N2=168 | × | × | × | N1=40, N2=10 |
| | | | Fold Change: 2.26 | | | | Fold Change: 3.04 |
| | | | P-value: 0 | | | | P-value: .00005 |
| 94 | 171 | AA584310 | 398.21+/-320.55 | | | 402.55+/-323.55 | 402.55+/-323.55 |
| | | | 1296.13+/-716.97 | | | 1058.09+/-583.59 | 1089.78+/-475.55 |
| | | | N1=40, N2=168 | × | × | N1=40, N2=31 | N1=40, N2=10 |
| | | | Fold Change: 3.84 | | | Fold Change: 3.12 | Fold Change: 3.31 |
| | | | P-value: 0 | | | P-value: 0 | P-value: .00032 |
| 95 | 172 | AA584403 | 593.26+/-1291.79 | | | 593.26+/-1291.79 | 593.26+/-1291.79 |
| | | | 85.5+/-209.13 | | | 113.63+/-199.33 | 77.07+/-76.22 |
| | | | N1=40, N2=168 | × | × | N1=40, N2=31 | N1=40, N2=10 |
| | | | Fold Change: 3.61 | | | Fold Change: 2.7 | Fold Change: 3.14 |
| | | | P-value: 0 | | | P-value: .00077 | P-value: .00412 |
| 8 | 174 | AA595800 | | | | 405.39+/-344.1 | |
| | | | | | | 167.2+/-159.75 | |
| | | | × | × | × | N1=40, N2=31 | × |
| | | | | | | Fold Change: 2.2 | |
| | | | | | | P-value: .00334 | |
| 46 | 176 | AA602521 | | | | | 258.64+/-109.45 |
| _ | | | | | | | 92.33+/-57.89 |
| | | | × | × | × | × | NI=39, N2=6 |
| | | | | | | | Fold Change: 3.01 |
| | | | | | | | P-value: .00015 |
| 86 | 178 | AA609310 | 289.22+/-160.55 | | | 285.39+/-160.8 | 285.39+/-160.8 |
| _ | | | 106.23+/-108.02 | | | 81.33+/-54.9 | 139.85+/-180.52 |
| | | | N1=40, N2=168 | × | × | N1=40, N2=31 | N1=40, N2=10 |
| | | | Fold Change: 3.16 | | | Fold Change: 3.57 | Fold Change: 2.65 |
| | | | P-value: 0 | | | P-value: 0 | P-value: .00931 |

| # | Sea TD | Genbank | Normal vs All | Normal vs Malignant | Normal vs Stage I | Normal vs Stage II | Normal vs Stage III |
|-----|--------|----------|-------------------|---------------------|-------------------|--------------------|---------------------|
| 8 | 180 | AA610522 | 803+/-768.74 | | 803+/-768.74 | 803+/-768.74 | |
| ` | 3 | | AF F31C-1+2C TC2C | | 1730 2+/-869 55 | 1878 88+/-1670.25 | |
| | | | N1=0 N2=168 | * | N1=40 N2=6 | N1=40 N2=31 | × |
| | | | 11 TO, 112 TO | • | Pold Changes 2 79 | Pold Change: 2 57 | |
| | | | P-value: 0 | | P-value: .00045 | P-value: .00552 | |
| 100 | 181 | AA612864 | | | | 228,36+/-116.88 | 228.36+/-116.88 |
| | | | | | | 495.29+/-290.05 | 440.68+/-155.22 |
| _ | | | × | × | × | N1=40, N2=31 | N1=40, N2=10 |
| _ | | | | | | Fold Change: 2.03 | Fold Change: 2.03 |
| | | | | | | P-value: 0 | P-value: .00004 |
| 101 | 184 | AA621478 | 394.08+/-322.25 | | 398.69+/-325.12 | 398,69+/-325.12 | 398.69+/-325.12 |
| | | | 60.41+/-84.83 | | 79.9+/-118.32 | 53,41+/-82.79 | 18+/-20.91 |
| | | | N1=40 N2=168 | × | N1=40, N2=6 | N1=40, N2=31 | N1=40, N2=10 |
| | | | Fold Change: 6.59 | | Fold Change: 5.51 | Fold Change: 7.48 | Fold Change: 11.4 |
| | | | P-value: 0 | | P-value: .00526 | P-value: 0 | P-value: 0 |
| 183 | 186 | AA625387 | | | | 92.36+/-35.34 | 92.36+/-35.34 |
| } | 1 | | | | | 208.73+/-105.36 | 231.14+/-178.29 |
| | | | × | × | × | N1=39, N2=31 | N1=39, N2=6 |
| | | | | | | Fold Change: 2.15 | Fold Change: 2.24 |
| | | | | | | P-value: 0 | P-value: .00162 |
| 103 | 180 | AA628467 | 1150.97+/-497.25 | | | 1145.06+/-502.33 | 1145.06+/-502.33 |
| _ | ì | | 290.19+/-235.1 | | | 274.87+/-259.13 | 122.9+/-92.36 |
| | | | N1=40, N2=168 | × | × | N1=40, N2=31 | N1=40, N2=10 |
| | | | Fold Change: 4.65 | | | Fold Change: 4.99 | Fold Change: 9.81 |
| _ | | | P-value: 0 | | | P-value: 0 | P-value: 0 |
| 101 | 192 | AA631215 | | | | 190,98+/-126,92 | |
| _ | | | | | | 541.33+/-898.12 | |
| | | | × | × | × | N1=40, N2=31 | × |
| | | | | | | Fold Change: 2.05 | |
| _ | | | | | | P-value: .00057 | |
| 105 | 193 | AA633203 | 127.67+/-220.56 | | | 130.61+/-222.66 | 130.61+/-222.66 |
| | | | 389.82+/-388.38 | | | 451.59+/-365.56 | 413.97+/-273.89 |
| | | | N1=40, N2=168 | × | × | N1=40, N2=31 | N1=40, N2=10 |
| | | | Fold Change: 3.74 | | | Fold Change: 5.08 | Fold Change: 4.59 |
| _ | | | P-value: 0 | | | P-vaine: 0 | P-value: .00196 |

| 46 | Sea ID | Genbank | Normal vs All | Normal vs Malignant | Normal vs Stage I | Normal vs Stage II | Normal vs Stage III |
|-----|--------|----------|----------------------|---------------------|-------------------|--------------------------------------|---------------------|
| 106 | 194 | AA634799 | 763,44+/-619.75 | | | 739.38+/-608.62 | |
| | | | 22 100 1.10 020 | | | 140 74±/-650 81 | |
| | | | 00.104 (N-11A | > | * | N1=40 N7=31 | × |
| | | | 10 TAT -40° TAT -10° | 4 | • | 7.11 | |
| | | | Fold Change: 2.41 | | | Food Change: 2.20 P-value: .00284 | |
| 107 | 196 | AA658561 | | | | | 238.89+/-114.83 |
| ì | , | | | | | | 110.57+/-54.76 |
| | | | × | × | × | × | N1=40, N2=10 |
| | | | ŗ | 1 | | | Fold Change: 2.02 |
| | | | | | | | P-value: .00612 |
| 108 | 198 | AA669106 | 82.724-129.9 | | | 84.29+/-131.22 | 84.29+/-131.22 |
| | | | 292.67+/-186.26 | | | 383.96+/-227.86 | 381.34+/-202.47 |
| | | | N1=40, N2=168 | × | × | N1=40, N2=31 | N1=40, N2=10 |
| | | | Fold Change: 4.65 | | | Fold Change: 6.02 | Fold Change: 6.04 |
| | | | P-value: 0 | | | P-value: 0 | P-value: 0 |
| 109 | 200 | AA700621 | 467.51+/-455.09 | | 467.51+/-455.09 | 467.51+/-455.09 | 467.51+/-455.09 |
| | | | 66.85+/-123.51 | | 87.68+/-141.47 | 36.61+/-66.8 | 26.41+/-35.67 |
| | | | N1=40, N2=168 | × | N1=40, N2=6 | N1=40, N2=31 | N1-40, N2-10 |
| | | | Fold Change: 6.01 | | Fold Change: 4.72 | Fold Change: 7.35 | Fold Change: 8.68 |
| | | | P-value: 0 | | P-value: .02358 | P-value: 0 | P-value: 0 |
| 110 | 204 | AA703262 | | | | 1213.89+/-750.28 | 1213.89+/-750.28 |
| | | | | | | 519.38+/-282.83 | 361.77+/-193.77 |
| | | | × | × | × | NI=40, NZ=31 | N1=40, N2=10 |
| | | | | | | Fold Change: 2.09 | Fold Change: 3.02 |
| | | | | | | P-value: .00044 | P-value: .00125 |
| 111 | 214 | AA742697 | 1009.7+/-1062.61 | | | 1026.03+/-1071.41 | 1026.03+/-1071.41 |
| | | | 222.47+/-326.55 | | | 238.53+/-383.92 | 142.42+/-134 |
| | | | N1=40, N2=168 | × | × | N1=40, N2=31 | N1=40, N2=10 |
| | | | Fold Change: 4.19 | | | Fold Change: 4.18 | Fold Change: 5.09 |
| | | | P-vaine: 0 | | | P-value: .00001 | P-value: .00012 |
| 112 | 215 | AA747315 | | | | 448.86+/-190.26 | 448.86+/-190.26 |
| | | | | | | 258.21+/-198.98 | 135.53+/-84.89 |
| | | | × | × | × | N1=40, N2=31 | N1=40, N2=10 |
| | | | | | | Fold Change: 2.02 | Fold Change: 3.66 |
| | | | | | | P-value: .00001 | P-value: .0001 |

| | | | | | TT | | |
|-----|--------|----------|-------------------|---------------------|-------------------|--------------------|--------------------------------------|
| * | Seq ID | Genbank | Normal vs All | Normal vs Malignant | Normal vs Stage I | Normal vs Stage II | Normal vs Stage III |
| 12 | 216 | AA760767 | 100.23+/-83.19 | | | 101.93+/-83.58 | 101.93+/-83.58 |
| 1 | 2 | | 345 71+1.246.25 | | | 273.79+/-236.49 | 388.04+/-242.5 |
| | | | M1-40 N2=168 | Þ | × | N1=40. N2=31 | N1=40. N2=10 |
| | | | Fold Change: 2 06 | • | | Fold Change: 2.59 | Fold Change: 3.64 |
| | | | P-value: 0 | | | P-value: .00001 | P-value: .00384 |
| 114 | 218 | AA772278 | | | | | 137.98+/-96.56 |
| | | | | | | | 296.31+/-168.39 |
| | | | × | × | × | × | N1=40, N2=10 |
| | | | | | | | Fold Change: 2.42 |
| | | | | | | | P-value: .00585 |
| Ľ. | 220 | AA775180 | | | | | 264.55+/-193.81 |
| 2 | | | | | | | 84.64+/-55.48 |
| | | | × | × | × | × | N1=40, N2=10 |
| | | | | | | | Fold Change: 2.66 |
| | | | | | | | P-value: .00029 |
| 116 | 223 | AA777369 | | | | 104.24+/-57.98 | |
| | | | | | | 211.86+/-128.62 | |
| | | | × | × | × | N1=40, N2=31 | × |
| | | | | | | Fold Change: 2.02 | |
| | | | | | | P-value: 0 | |
| 117 | 226 | AA779795 | | | | | 493.39+/-168.5 |
| | | | | 1 | ; | , | 21.16.16.15.15. |
| | | | × | × | × | < | NI=#0, NZ=10 |
| | | | | | | | Fold Change: 2.25 P-value: .00072 |
| 118 | 227 | AA788946 | 568.73+/-336.98 | | | | |
| ŀ | | | 1313.01+/-867.03 | | | | |
| | | | N1=39, N2=168 | × | × | × | × |
| | | | Fold Change: 2.15 | | | | |
| | | | P-value: 0 | | | | |
| 119 | 228 | AA789332 | 120.94+/-61.86 | | | 120.94+/-61.86 | |
| | | | 221.57+/-96.99 | | | 228.17+/-103.03 | 1 |
| | | | N1=39, N2=168 | × | × | N1=39, N2=31 | × |
| | | | Fold Change: 1.91 | | | Fold Change: 1.94 | |
| | | | P-value: 0 | | | P-value: .00007 | |

| 78 | Seq ID | Genbank | Normal vs All | Normal vs Malignant | Normal vs Stage I | Normal vs Stage II | Normal vs Stage III |
|-----|--------|--|-------------------|--|-------------------|--------------------|---------------------|
| 120 | 230 | AA807154 | 271.93+/-107.94 | | | 273.39+/-108.95 | 273.39+/-108.95 |
| | | | 150.18+/-94.12 | | | 115.85+/-82.09 | 137.82+/-101.26 |
| | | | N1-40, N2=168 | × | × | N1=40, N2=31 | N1=40, N2=10 |
| | | | Fold Change: 2.01 | | | Fold Change: 2.55 | Fold Change: 2.3 |
| | | | P-value: 0 | | | P-value: 0 | P-value: .00953 |
| 121 | 235 | AA830844 | 93.3+/-120.3 | | | 94.7+/-121.54 | 94.7+/-121.54 |
| | | | 272.03+/-321.32 | | | 320.78+/-455.06 | 361.79+/-369.37 |
| | | | N1=40, N2=168 | × | × | N1=40, N2=31 | N1=40, N2=10 |
| | | | Fold Change: 2.64 | ֖֓֞֟֝֓֓֓֓֓֓֓֓֓֓֓֓֓֓֓֓֓֓֓֓֓֓֓֓֓֓֓֓֓֓֓֓֓֓֓ | | Fold Change: 2.9 | Fold Change: 3.88 |
| | | | P-value: 0 | | | P-value: .00003 | P-value: .00167 |
| 122 | 236 | AA838843 | | | | | 1133,4+/-889.08 |
| | | | | | • | | 503.96+/-209.26 |
| | | | × | × | × | × | N1=40, N2=10 |
| | | | | | | | Fold Change: 2 |
| | | | | | | | P-value: .00069 |
| 123 | 239 | AA846091 | 83,83+/-80,09 | | 84.12+/-81.11 | 84.12+/-81.11 | |
| | | | 254,24+/-227,12 | | 271.23+/-192.47 | 226.57+/-179.06 | |
| | | | N1=40, N2=168 | × | N1=40, N2=6 | N1=40, N2=31 | × |
| | | | Fold Change: 2,88 | | Fold Change: 3.44 | Fold Change: 2.75 | |
| | | | P-value: 0 | | P-value: .01151 | P-value: 0 | |
| 124 | 241 | A A 875998 | | | | | 144.49+/-132.98 |
| • | ŧ | OCCUPATION AND ADDRESS OF THE PARTY OF THE P | | | | | 225.7+/-97.08 |
| | | | × | × | × | × | N1=40, N2=10 |
| | | | | | | | Fold Change: 2.02 |
| | | | | | | | P-value: .00162 |
| 125 | 246 | AA909042 | | | | 166.1+/-92.52 | 166.1+/-92.52 |
| | | | | | | 345.62+/-200.16 | 375,51+/-207,38 |
| | | | × | × | × | N1=40, N2=31 | N1=40, N2=10 |
| | | | | | | Fold Change: 2.07 | Fold Change: 2.2 |
| | | | | | | P-value: 0 | P-value: .00481 |
| 126 | 249 | 826606VV | | | | 304.09+/-148.66 | |
| | | | | | | 159.71+/-101.37 | |
| | | | × | × | × | N1=40, N2=31 | × |
| | | | | | | Fold Change: 2.04 | |
| | | | ! | | | P-value: .00003 | |

| | | | | | A management of the second sec | | |
|-----|--------|------------|-------------------|---------------------|--|--------------------|--------------------------------|
| # | Sed ID | Genbank | Normal vs All | Normal vs Malignant | Normal vs Stage I | Normal vs Stage II | Normal vs Stage III |
| 127 | 251 | AA913079 | | | | | 379.55+/-192 819.72+/-391.8 |
| | | | × | × | × | × | N1=40, N2=10 |
| | | | | | | | Fold Change: 2.18 |
| | | | | | | | P-value: .00102 |
| 128 | 254 | AA921830 | 91.7+/-113.88 | | | 92,93+/-115.1 | 92,93+/-115.1 |
| | | | 226.43+/-153.69 | | | 257,36+/-191.81 | 324.58+/-186.68 |
| | | | N1=40, N2=168 | × | × | N1=40, N2=31 | N1=40, N2=10 |
| _ | | | Fold Change: 2.9 | | | Fold Change: 3.24 | Fold Change: 4 |
| | | | P-value: 0 | | | P-value: 0 | P-value: .0002 |
| 129 | 255 | AA921922 | 312,62+/-288.86 | | | 312,44+/-292.63 | 312.44+/-292.63 |
| | | | 76.08+/-72.69 | | | 75.75+/-54.61 | 36.4+/-15.92 |
| | | | N1=40, N2=168 | × | × | N1=40, N2=31 | N1=40, N2=10 |
| | | | Fold Change: 4.16 | | | Fold Change: 3.86 | Fold Change: 6.83 |
| | | | P-value: 0 | | | P-value: 0 | P-value: 0 |
| 130 | 257 | AA928698 | | | | | 979.86+/-357.7 |
| | | | | | | | 383.02+/-284.05 |
| | | | × | × | × | × | N1=40, N2=10 |
| _ | | | | | | | Fold Change: 3.22 |
| | | | | | | | P-value: .00199 |
| 13 | 258 | A A 928776 | 334.78+/-156.59 | | | 334.78+/-156.59 | 334.78+/-156.59 |
| | | | 126.01+/-82.79 | | | 129.6+/-115.01 | 91.05+/-43.12 |
| _ | | | N1=39, N2=168 | × | × | N1=39, N2=31 | N1=39, N2=6 |
| _ | | | Fold Change: 2.72 | | | Fold Change: 2.74 | Fold Change: 3.62 |
| _ | | | P-value: 0 | | | P-value: 0 | P-value: .00002 |
| 132 | 259 | AA928876 | | | | 279.93+/-242.15 | |
| _ | | | × | × | × | N1=40, N2=31 | × |
| | | | • | : | | Fold Change: 2.37 | |
| | | | | | | P-value: .00023 | |
| 133 | 260 | AA936632 | 124.77+/-125.66 | | | 125.03+/-127.3 | 125.03+/-127.3 |
| | | | 335.73+/-304.1 | | | 307.81+/-176.76 | 325.44+/-181.42 |
| | | | N1=40, N2=168 | × | × | N1=40, N2=31 | N1-40, N2=10 |
| | | | Fold Change: 2.65 | | | Fold Change: 2.68 | Fold Change: 3 |
| | | | P-value: 0 | | | L-Varies o | Y Adding Conces |

| * | Sed ID | Genbank | Normal vs All | Normal vs Malignant | Normal vs Stage I | Normal vs Stage II | Normal vs Stage III |
|-----|--------|------------|-------------------|---------------------|-------------------|--|---------------------|
| 134 | 261 | AA947123 | 288.72+/-99.04 | | | 287.11+/-99.81 | 287.11+/-99.81 |
| | | | 125 0647 76 03 | | | 100 201/ 6/ 69 | 25 72 /407 901 |
| | | | 20.01=1.00.051 | 24 | þ | 10 01 01 11 11 11 11 11 11 11 11 11 11 1 | 21 CT 1 CT 1 CT 1 |
| | | | NI-40, INZ-106 | < | < | IN 1-40, INZ-31 | NI-+0, 142-10 |
| | | | Fold Change: 2.34 | | | Fold Change: 2.88 | Fold Change: 3.03 |
| | | | P-value: 0 | | | P-value: 0 | P-value: .00116 |
| 135 | 263 | AA968657 | 120.64+/-123.71 | | | | |
| | | | 221.47+/-188.53 | | | | |
| | | | N1=40 N2=168 | Α | >- | > | > |
| | | | Fold Change: 2 02 | < | < | < | < |
| | | | P-value: ,00055 | | | | |
| 136 | 264 | A A 969863 | | | | | 154.92+/-62.69 |
| ì | | | | | | | 335+/-158.01 |
| | | | Þ | Þ | Þ | Þ | NT-40 NT-10 |
| | | | < | 4 | < | < | OI-7NI OF IN |
| | | | | | | | Fold Change: 2.11 |
| | | | | | | | r-value: .00036 |
| 137 | 506 | AA976064 | 363.92+/-151.16 | | | | |
| | | | 183.7+7-101.29 | | | | |
| | | | N1=40, N2=168 | × | × | × | × |
| | | | Fold Change: 2.17 | | | | |
| | | | P-value: 0 | | | | |
| 138 | 292 | AA977204 | | | | | 159.24+/-78.21 |
| | | | | | | | 300 74+/-74 19 |
| | | | 4 | þ | Þ | Þ | A11-40 A10-10 |
| | | | * | * | * | × | NI=40, NZ=10 |
| | | | | | | | Fold Change: 2.02 |
| | | | | | | | P-value: 0 |
| 139 | 272 | AB007972 | 317.37+/-102.68 | | | 317.37+/-102.68 | 317.37+/-102.68 |
| | | | 141.41+/-56.37 | | | 127.42+/-50.63 | 110.2+/-25.69 |
| | | | N1=40, N2=168 | × | × | N1=40, N2=31 | N1-40, N2-10 |
| | | | Fold Change: 2.26 | | | Fold Change: 2.49 | Fold Change: 2.75 |
| | | | P-value: 0 | | | P-value: 0 | P-value: 0 |
| 140 | 280 | AC004010 | | | | 237.41+/-101.03 | |
| | | | | | | 119.13+/-50.91 | |
| | | | × | × | × | N1=39, N2=31 | × |
| | | | | | | Fold Change: 2.01 | |
| | | | | | | Pavalue: 0 | |

| * | Seq ID | Genbank | Normal vs All | Normal vs Malignant | Normal vs Stage I | Normal vs Stage II | Normal vs Stage III |
|-----|--------|----------|-------------------|---------------------|-------------------|--------------------|---------------------|
| 4 | 281 | AC004770 | 116.54+/-76.22 | | | 116.54+/-76.22 | 116.54+/-76.22 |
| | | | 231.2+/-112.79 | | | 258.92+/-95.73 | 353.08+/-90.31 |
| | | | N1=40, N2=168 | × | × | N1=40, N2=31 | N1=40, N2=10 |
| | | | Fold Change: 2.03 | | | Fold Change: 2.41 | Fold Change: 3.44 |
| | | | P-value: 0 | | | P-value: 0 | P-value: 0 |
| 142 | 286 | AF007150 | | | | | 560.67+/-252.85 |
| | | | | | | | 244.22+/-118.81 |
| | | | × | × | × | × | NI=40, N2=10 |
| | | | | | | | Fold Change: 2.3 |
| | | | | | | | P-value: .00021 |
| 143 | 287 | AF009314 | 280.81+/-113.17 | | | | 280.81+/-113.17 |
| | | | 138,34+/-66.55 | | | | 90,09+/-43.95 |
| | | | N1=40, N2=168 | × | × | × | N1=40, N2=10 |
| | | | Fold Change: 2.04 | | | | Fold Change: 3.18 |
| | | | P-value: 0 | | | | P-value: .00003 |
| 144 | 297 | AF052142 | 307.17+/-169.55 | | | 307.17+/-169.55 | 307.17+/-169.55 |
| | | | 136.07+/-86.42 | | | 134.05+/-84.74 | 131.75+/-87.96 |
| | | | N1=40, N2=168 | × | × | N1=40, N2=31 | N1=40, N2=10 |
| | | | Fold Change: 2.28 | | | Fold Change: 2.25 | Fold Change: 2.44 |
| | | | P-value: 0 | | | P-value: 0 | P-value: .00309 |
| 145 | 300 | AF058075 | | | | | -15.62+/-107.9 |
| | | | | | | | 355,75+/-489,88 |
| | | | × | × | × | × | N1=40, N2=10 |
| | | | | | | | Fold Change: 3.73 |
| | | | | | | | P-value: .0394 |
| 146 | 304 | AF070648 | 1031,52+/-429,52 | | | 1031.52+/-429.52 | 1031.52+/-429.52 |
| | | | 432.03+/-213.34 | | | 357.01+/-152.1 | 292.72+/-163.49 |
| | | | N1=40, N2=168 | × | × | NI=40, N2=31 | N1=40, N2=10 |
| | | | Fold Change: 2.41 | | | Fold Change: 2.82 | Fold Change: 3.62 |
| | | | P-value: 0 | | | P-value: 0 | P-value: .00002 |
| 147 | 300 | AF150174 | 211.02+/-104.05 | | | 211.02+/-104.05 | 211.02+/-104.05 |
| | | | 86.63+/-46.76 | | | 86.92+/-51.98 | 78.32+/-31.52 |
| | | | N1=39, N2=168 | × | × | N1=39, N2=31 | N1=39, N2=6 |
| | | | Fold Change: 2.26 | | | Fold Change: 2.26 | Fold Change: 2.38 |
| | | | P-value: 0 | | | P-value: 0 | P-value: 0 |

| # | Sed ID | Genbank | Normal vs All | Normal vs Malignant | Normal vs Stage I | Normal vs Stage II | Normal vs Stage III |
|-----|--------|----------|-------------------|---------------------|-------------------|--------------------|---------------------|
| 148 | 311 | AI002238 | | | | | 254.67+/-91.81 |
| | | | × | × | × | × | N1=40, N2=10 |
| | | | | | | | Fold Change: 2.72 |
| | | | | | | | P-value: .00083 |
| 149 | 314 | AI016604 | | | | 502,7+/-255.95 | 502.7+/-255.95 |
| | , | | | | | 262.05+/-145.41 | 226.05+/-112.74 |
| | | | × | × | × | N1=40, N2=31 | N1=40, N2=10 |
| | | | | | | Fold Change: 2.02 | Fold Change: 2.18 |
| | | | | | | P-value: .00002 | P-value: .00111 |
| 120 | 317 | AI018523 | 422.08+/-187.64 | | | 422.08+/-187.64 | 422.08+/-187.64 |
| | | | 94,36+/-123.53 | | | 77.78+/-105.86 | 44.08+/-59.8 |
| | | | N1=40, N2=168 | × | × | N1=40, N2=31 | N1=40, N2=10 |
| | | | Fold Change: 5.56 | | | Fold Change: 6.54 | Fold Change: 9.77 |
| | | | P-value: 0 | | | P-value: 0 | P-value: 0 |
| 151 | 321 | AI031771 | 85.9+/-105.07 | | | 85.9+/-105.07 | |
| | | | 257.24+/-405.17 | | | 298.84+/-697.75 | |
| | | | N1=40, N2=168 | × | × | N1=40, N2=31 | × |
| | | | Fold Change: 2.53 | | | Fold Change: 2.33 | |
| | | | P-value: 0 | | | P-value: .00142 | |
| 152 | 324 | AI039005 | 201.84+/-130.44 | | | 203.54+/-131.69 | 203.54+/-131.69 |
| | | | 77.34+/-61.5 | | | 61.56+/-42.38 | 44.43+/-54.48 |
| | | | N1=40, N2=168 | × | × | N1=40, N2=31 | N1=40, N2=10 |
| | | | Fold Change: 2.58 | | | Fold Change: 3.03 | Fold Change: 3.97 |
| | | | P-value: 0 | | | P-value: 0 | P-value: .0001 |
| 153 | 325 | AI039722 | 990.63+/-1152.38 | | | 1007.24+/-1162.59 | 1007.24+/-1162.59 |
| | | | 438.37+/-1004.04 | | | 232.49+/-466.94 | 207.73+/-245.94 |
| | | | N1=40, N2=168 | × | × | N1=40, N2=31 | N1=40, N2=10 |
| | | | Fold Change: 3.87 | | | Fold Change: 6.17 | Fold Change: 5.76 |
| | | | P-value: 0 | | | P-value: 0 | P-value: .00392 |
| 154 | 326 | AI049549 | 653.84+/-535.62 | | | 653.84+/-535.62 | 653.84+/-535.62 |
| | | | 267.97+/-227.53 | | | 226.29+/-141.74 | 155.78+/-116.9 |
| | | | N1=40, N2=168 | × | × | N1=40, N2=31 | N1=40, N2=10 |
| | | | Fold Change: 2.36 | | | Fold Change: 2.5 | Fold Change: 3.66 |
| | | | P-value: 0 | | | P-value: .00001 | P-value: .00006 |

| | Seq ID | Genbank | Normal vs All | Normal vs Malignant | Normal vs Stage I | Normal vs Stage II | Normal vs Stage III |
|-----|--------|----------|-------------------|---------------------|--------------------------------|--------------------|---------------------|
| | 327 | AI049973 | | | | | 391.85+/-108.68 |
| | | | × | × | × | × | N1=39, N2=6 |
| | | | 1 | 1 | | | Fold Change: 1.95 |
| | | | | | | | P-value: .00002 |
| 156 | 330 | AI056241 | 241.39+/-152.04 | | 241.39+/-152.04 | 241.39+/-152.04 | 241.39+/-152.04 |
| | | | 67.68+/-55.35 | | 93.33+/-104.89 | 49.94+/-43.13 | 45.49+/-48.07 |
| | | | N1=39, N2=168 | × | N1=39, N2=10 | N1=39, N2=31 | N1=39, N2=6 |
| | | | Fold Change: 2.88 | | Fold Change: 2.9 | Fold Change: 3.33 | Fold Change: 4.03 |
| | | | P-value: 0 | | P-value: .03101 | P-value: 0 | P-value: .00003 |
| | 331 | AI057450 | 381.32+/-1572.07 | | | 381,32+/-1572.07 | 381.32+/-1572.07 |
| | | | 20.72+/-180.68 | | | 27.54-176.72 | 10.16+/-47.15 |
| | | | N1=40, N2=168 | × | × | N1=40, N2=31 | N1=40, N2=10 |
| | | | Fold Change: 2.92 | | | Fold Change: 2.87 | Fold Change: 2.99 |
| | | | P-value: .00006 | | | P-value: .00026 | P-value: .00095 |
| 158 | 332 | AI057637 | 568.22+/-502.47 | | 573.524/-507.9 | 573.52+/-507.9 | 573.52+/-507.9 |
| | | | 153,6+/-81.89 | | 163.69+/-59.04 | 129.31+/-59.91 | 124.36+/-39.47 |
| | | | N1=40, N2=168 | × | N1=40, N2=6 | N1=40, N2=31 | N1=40, N2=10 |
| | | | Fold Change: 3.29 | | Fold Change: 2.9 | Fold Change: 3.79 | Fold Change: 3.81 |
| | | | P-value: 0 | | P-value: .0001 | P-value: 0 | P-value: 0 |
| 159 | 336 | AI078033 | 454.77+/-280.5 | | | 454.77+/-280.5 | 454.77+/-280.5 |
| | | | 203.08+/-158.36 | | | 183.47+/-128.31 | 129.56+/-78.11 |
| | | | N1=40, N2=168 | × | × | N1=40, N2=31 | N1=40, N2=10 |
| | | | Fold Change: 2.27 | | | Fold Change: 2.42 | Fold Change: 3.22 |
| | | | P-value: 0 | | | P-value: 0 | P-value: .00008 |
| | 337 | AI078121 | 267.92+/-121.01 | | | | 267.92+/-121.01 |
| | | | 129.6+/-75.04 | | | | 124.52+/-39.11 |
| | | | N1=39, N2=168 | × | × | × | N1=39, N2=6 |
| | | | Fold Change: 2.01 | | | | Fold Change: 2.05 |
| | | | P-value: 0 | | | | P-value: .00004 |
| | 338 | AI079545 | | | 248.94+/-138.38 | | |
| | | | × | × | 465.5947-104.61 N1=40, N2=6 | × | × |
| | | | ! | | Fold Change: 2.14 | | |
| | | | | | P-value: .00004 | | |

| # | Sea TD | Genbank | Normal vs All | Normal vs Malignant | Normal vs Stage I | Normal vs Stage II | Normal vs Stage III |
|-----|--------|-------------------|--|---------------------|---|--|--|
| 162 | 339 | AI079558 | × | × | × | 115.76+/-141.71 214.96+/-122.84 N1=40, N2=31 Fold Change: 2.3 P-value: .00001 | × |
| 163 | 341 | A1083598 | 344.81+/-287.52 151.49+/-429.97 N1=40, N2=168 Fold Change: 3.75 P-value: 0 | × | × | 339.56+/-289.33 210.77+/-736.98 N1=40, N2=31 Fold Change: 4.41 P-value: 0 | 339.56+/-289.33 31.3+/-28.36 N1=40, N2=10 Fold Change: 6.76 P-yalue: 0 |
| 164 | 342 | AI08 66 14 | 300.02+/-151.08 127.26+/-88.54 N1=40, N2=168 Fold Change: 2.5 P-value: 0 | × | × | 301.2+/-152.86 96.79+/-68.03 N1=40, N2=31 Fold Change: 3.25 P-value: 0 | 301.2+/-152.86 68.08+/-44.9 N1=40, N2=10 Fold Change: 3.7 P-value: 0 |
| 165 | 343 | AI087975 | 137.52+/-92.28 277.44+/-204.79 N1=39, N2=168 Fold Change: 1.96 P-value: 0 | × | × | × | × |
| 166 | 344 | AI088609 | 711.92+/-592.71 327.21+/-706.4 N1=40, N2=168 Pold Change: 3.75 P-value: 0 | × | × | 709.25+/-600.21 163.7+/-218.98 N1=40, N2=31 Fold Change: 4.73 P-value: 0 | 709.25+/-600.21 78.97+/-52.81 N1=40, N2=10 Fold Change: 6.93 P-value: 0 |
| 167 | 345 | AI091154 | 351.29+/-406.17 73.95+/-133.05 N1=40, N2=168 Fold Change: 4.29 P-value: 0 | X | 351.29+/-406.17 87.21+/-103.53 N1=40, N2=6 Fold Change: 3.04 P-value: .0365 | 351.29+/-406.17 86.99+/-153.12 N1=40, N2=31 Fold Change: 3.61 P-value: 0 | 351.29+/-406.17 62.86+/-69.84 N1=40, N2=10 Fold Change: 3.91 P-value: .00107 |
| 168 | 346 | AI092936 | 185.51+/-150.9 417.08+/-227.5 N1=40, N2=168 Fold Change: 2.2 P-value: 0 | x | × | 186.98+/-152.58 425.95+/-204.95 N1=40, N2=31 Fold Change: 2.35 P-value: .00001 | × |

| Normal vs Stage III | | × | X 354.78+1.131.25 184.82+1.126.35 N1=40, N2=10 Pold Change: 2.08 Powthe: .00216 | X 354.78+/131.25 184.82+/126.35 Nather, 102216 500+/154.6 500+/154.6 500+/154.6 Nather, 102216 Nather, 102216 Nather, 102216 Soots-154.6 S | X 354.784/131.25 164.824-126.35 Nod Change: 2.08 Rold Change: 2.08 Rold Change: 2.08 Nod-1.0016 300+1.160.30 Nod-1.0016 Rold Change: 4.97 Pende Change: 4.97 Pende Change: 4.97 Pende Change: 2.66 Rold Change: 2.66 Rold Change: 2.66 Rold Change: 2.66 Rold Change: 2.46 Rold Change: 2.47 Rold Change: 2.46 Rold Change: 2.47 Rold Change: 2. | X 354 784-131.25 194 284-131.25 194 284-135.37 194 284-135.37 195 284-135.37 195 284-135.37 195 284-135.39 195 284-135.39 195 284-135.39 195 284-135.39 195 284-135.39 195 284-39.39 195 284-39.39 195 284-39.39 195 284-39.39 195 284-39.39 X | X 344784-131.25 184.824-126.35 184.824-126.35 184.824-126.35 184.824-126.35 184.824-126.35 184.824-126.39 184.824-126.39 185.844-126.39 185.84-142.18 185.84 |
|---|------------------|-------------|--|--|--|---|--|
| | × | | 354.78+/- 184.82+/- N1=40, N Fold Chang P-value. | | | | |
| × | | | | | | | |
| × | | × | | 300+/-164.6 67.15+/-70.68 N1=40, N2=31 Fold Change: 4.66 P-value: 0 | 300+/164.6 67.15+/-70.68 NI=40, N2=31 NI=40, N2=31 Portlue: 0 297.764/-110.04 152.584/-82.32 NI=39, N2=31 Podd Change: 2.04 P-value: 0 | 300+/164.6 (7.15+/-70.68 N1=40, N2=21 Fold Changes 4.66 P-value: 0 | 300+/164.6 300+/164.6 N1-40, N2-21 Pold Change: 4,66 Pold Change: 4,66 Pold Change: 4,67 157.548+120.2 Pold Change: 2,07 |
| * ^ | ^ | | 300+/- | | | | |
| × × | × | | +/-164.6 12+/-67 | 40, N2=6 hange: 3.22 ue: .00482 | 40, N2=6 thange: 3.22 uc: .00482 X | Ado, N2=6 thange: 3.22 ue: .00482 X | A X X X X X X X X X X X X X X X X X X X |
| X X X 300+/-164.6 | X 300+/-164.6 | 300+/-164.6 | 91.12+/-67 N1=40, N2=6 | Fold Change: 3.22 P-value: .00482 | old Change: 3.7 P-value: .0048. | old Changer 3. P-value: .00482 X X X | old Changer 3. P-value: ,0048; X X X |
| 30 | 300 | 300 | N S | P-va | P-va | F-va | Poy |
| | | | | | | , | * |
| * × | × | | × | | × | × × | × × × |
| | | | | | | | |
| 244.3847-121.07 348.3847-121.07 N1=39, N2=168 Fold Change: 1.75 P-value: .00003 | | × | 300.32+/-162.49 75.22+/-66.82 | N1=40, N2=168 Fold Change: 4.19 P-value: 0 | N1=40, N2=168 Fold Change: 4.19 P-value: 0 297.76+4-110.04 156.35+7-127.73 N1=39, N2=168 Fold Change: 2.06 P-value: 0 P-value: 0 | NI=40, N2=168 Povalue: 0 Povalue: 0 Povalue: 0 156.254-127.73 NI=29, N2=168 Povalue: 0 Povalue: 0 Povalue: 0 Povalue: 0 Povalue: 0 Povalue: 0 NI=29, N2=168 NI=29, N2=168 NI=29, N2=168 NI=29, N2=168 NI=29, N2=168 Povalue: 0 | Fold Changes 4.19 Fold Changes 4.19 Fold Changes 4.19 Fold Changes 1.10 Fordhold 116.534+1.277.73 Fordhold Changes 2.06 Fold Changes 2.06 Fold Changes 2.06 Fold Changes 2.08 Fold Changes 2.28 Fold Changes 2.31 |
| A1094535 | | AI096389 | AT123555 | | AII23738 | AI123738 AI125252 | AI123738 AI125252 AI126237 |
| | | ľ | | | | | |
| 248 | | 349 | 351 | | 352 | i i | |
| | | 0.1 | | 1_ | 1 2 | n n | 171 173 173 |

| # | Seq ID | Genbank | Normal vs All | Normal vs Malignant | Normal vs Stage I | Normal vs Stage II | Normal vs Stage III |
|-----|--------|----------|-------------------|---------------------|-------------------|--------------------|---------------------|
| 176 | 359 | AI128820 | 225.63+/-90.11 | | | 224.42+/-90.96 | 224.42+/-90.96 |
| | | | 101.84+/-66.82 | | | 96.42+/-39.89 | 82.27+/-38.37 |
| | | | N1=40, N2=168 | × | × | N1=40, N2=31 | N1=40, N2=10 |
| _ | | | Fold Change: 2.29 | | | Fold Change: 2.29 | Fold Change: 2.85 |
| _ | | | P-value: 0 | | | P-value: 0 | P-value: .0008 |
| 177 | 360 | AI129320 | 462.88+/-248.98 | | | 462.06+/-252.18 | 462.06+/-252.18 |
| | | | 223.69+/-111.28 | | | 218.45+/-156.49 | 242.37+/-237.11 |
| | | | N1=40, N2=168 | × | × | N1=40, N2=31 | N1=40, N2=10 |
| | | | Fold Change: 2.08 | | | Fold Change: 2.3 | Fold Change: 2.23 |
| | | | P-value: 0 | | | P-value: 0 | P-value: .00458 |
| 178 | 362 | AI131078 | 299.48+/-223.81 | | | 299.48+/-223.81 | 299.48+/-223.81 |
| | | | 105.31+/-56.6 | | | 102.41+/-57.3 | 99.18+/-37.7 |
| | | | N1=40, N2=168 | × | × | N1=40, N2=31 | N1=40, N2=10 |
| | | | Fold Change: 2.51 | | | Fold Change: 2.6 | Fold Change: 2.48 |
| _ | | | P-value: 0 | | | P-value: 0 | P-value: .00001 |
| 179 | 365 | AI140764 | 241.9+/-66.27 | | | | 241.9+/-66.27 |
| | | | 127.44+/-57.54 | | | | 119.16+/-48.86 |
| | | | N1-39, N2-168 | × | × | × | N1=39, N2=6 |
| _ | | | Fold Change: 2.03 | | | | Fold Change: 2.11 |
| | | | P-value: 0 | | | | P-value: .00026 |
| 180 | 298 | AT141556 | | | | | 297.58+/-90.49 |
| | į | | | | | | 101.69+/-39.39 |
| | | | × | × | × | × | N1=40, N2=10 |
| | | | | | | | Fold Change: 2.97 |
| | | | | | | | P-value: 0 |
| 181 | 368 | AI144477 | 560.95+/-250.34 | | | 564.89+/-252.35 | |
| | | | 250.3+/-192.18 | | | 332.2+/-275.56 | _ |
| | | | N1=40, N2=168 | × | × | N1=40, N2=31 | × |
| _ | | | Fold Change: 2.61 | | | Fold Change: 2.04 | |
| | | | P-value: 0 | | | P-value: .00003 | |
| 182 | 370 | AI148006 | 241.32+/-191.01 | | | 241.17+/-193.5 | 241.17+/-193.5 |
| | | | 84.05+/-116.71 | | | 51.87+/-67.91 | 25.24+/-42.83 |
| | | | N1=40, N2=168 | × | × | N1=40, N2=31 | N1=40, N2=10 |
| | | | Fold Change: 2.99 | | | Fold Change: 4.03 | Fold Change: 5.56 |
| | | | P-value: 0 | | | P-value: 0 | P-value: 0 |

| | 3 | | | _ | | | ż | | | | | | | | | _ | _ | | _ | _ | _ | | _ | | | | _ | | _ | |
|---------------------------------------|-----------------|---------------|--------------------------------------|--------------|--------------|-------------------|----------------|---------------|---------------|-------------------|-----------------|----------------|----------------|------------------------------------|------------|-----------------|----------------|--------------|-------------------|-----------------|----------------|----------------|--------------|------------------|-----------------|-----------------|----------------|--------------|-------------------|-----------------|
| Normal vs Stage III | .223.18+/:91.72 | N1=39, N2=6 | Fold Change: 2.37 P-value: .00396 | 243+/-112.73 | N1=40, N2=10 | Fold Change: 2.04 | 301.58+/-452.1 | 76.68+/-82.57 | N1=40, N2=10 | Fold Change: 2.77 | P-value: .00751 | | | × | | . 284.7+/-101.6 | 127.96+/-60.39 | N1=40, N2=10 | Fold Change: 2.41 | P-value: .00278 | 205.71+/481.06 | 101.78+/-59.91 | N1=40, N2=10 | Fold Change: 2.2 | P-value: .00676 | 290.63+/-103.17 | 100.98+/-38.82 | N1=40, N2=10 | Fold Change: 2.9 | P-value: .00001 |
| Normal vs Stage II | 223.18+/-91.72 | N1=39, N2=31 | Fold Change: 2.25 P-value: 0 | | × | | 301.58+/-452.1 | 76.69+/-52.21 | N1=40, N2=31 | Fold Change: 2.46 | P-value: .00013 | | | × | | | | × | | | | | × | | | 290.63+/-103.17 | 141.14+/-64.83 | N1=40, N2=31 | Fold Change: 2.13 | P-value: 0 |
| Normal vs Stage I | 7 | × | | | × | | | | × | | | | | × | | | | × | | | | | × | | | | | × | | |
| Normal vs Malignant Normal vs Stage I | | × | | | × | | | | × | | | | 1 | × | | | | × | | | | | × | | | | | × | | |
| Normal vs All | 223.18+/-91.72 | N1=39, N2=168 | Fold Change: 2.5 P-value: 0 | | × | | 301.58+/-452.1 | 77.97+/-71.53 | N1=40, N2=168 | Fold Change: 2.61 | P-value: .00001 | 356.99+/-119.9 | 188.39+/-84.68 | N1=39, N2=168 Fold Change: 1.94 | P-value: 0 | | | × | | | | | × | | | | | × | | |
| Genbank | AI149693 | | | AI160811 | | | AI161049 | | | | | AI168057 | | | | AI189011 | | | | | AI189255 | | | | | AI190755 | | | | |
| Seq ID | 373 | | | 375 | | | 376 | | | | | 378 | | | | 380 | | | | | 381 | | | | | 382 | | | | |
| * | 183 | | | 184 | | | 185 | _ | | | | 186 | | | | 187 | | | | | 188 | _ | | _ | | 189 | | _ | | |

| # | Sed ID | Genbank | Normal vs All | Normal vs Malignant | Normal vs Stage I | Normal vs Stage II | Normal vs Stage III |
|-----|--------|----------|-------------------|---------------------|-------------------|--------------------|---------------------|
| 190 | 384 | AI200954 | 529.23+/-316.46 | | | 524.84+/-319.36 | 524,84+/-319,36 |
| | | | 256.1+/-158.56 | | | 219.57+/-119.93 | 165.98+/-93.19 |
| | | | N1=40, N2=168 | × | × | N1=40, N2=31 | N1=40, N2=10 |
| | | | Fold Change: 2.07 | | | Fold Change: 2.28 | Fold Change: 3.18 |
| | | | P-value: 0 | | | P-value: .00001 | P-value: .00195 |
| 161 | 385 | AI201273 | 535.25+/-296.2 | | | 533.25+/-299.79 | 533.25+/-299.79 |
| | | | 238.85+/-188.24 | | | 203.14+/-125.12 | 125.11+/-95.45 |
| | | | N1=40, N2=168 | × | × | N1=40, N2=31 | N1=40, N2=10 |
| | | | Fold Change: 2.5 | | | Fold Change: 2.61 | Fold Change: 4.56 |
| | | | P-value: 0 | | | P-value: 0 | P-value: .0001 |
| 192 | 386 | AI201965 | 237.22+/-148.65 | | | 234.24+/-149.37 | 234.24+/-149.37 |
| | | | 107.53+/-70.7 | | | 100.47+/-56.35 | 65.63+/-51.87 |
| | | | N1=40, N2=168 | × | × | N1=40, N2=31 | NI=40, N2=10 |
| | | | Fold Change: 2.18 | | | Fold Change: 2.19 | Fold Change: 3.63 |
| | | | P-value: 0 | | | P-value: .00001 | P-value: .00031 |
| 193 | 387 | AT201982 | 279.97+/-146.77 | | | 279.97+/-146.77 | 279.97+/-146.77 |
| | | | 114,06+/-74.45 | | | 121,58+/-55,99 | 121.2+/-81.62 |
| | | | N1=39, N2=168 | × | × | N1=39, N2=31 | N1=39, N2=6 |
| | | | Fold Change: 2.45 | | | Fold Change: 2.14 | Fold Change: 2.51 |
| | | | P-value: 0 | | | P-value: 0 | P-value: .00327 |
| 194 | 388 | AI206014 | 213.72+/-107.69 | | | 211.4+/-108.08 | 211.4+/-108.08 |
| i | | | 81.38+/-54.34 | | | 80.55+/-42.41 | 60.05+/-31.32 |
| | | | N1=40, N2=168 | × | × | N1=40, N2=31 | N1=40, N2=10 |
| | | | Fold Change: 2.63 | | | Fold Change: 2.54 | Fold Change: 3.36 |
| | | | P-value: 0 | | | P-value: 0 | P-value: .00001 |
| 195 | 389 | AT206063 | 133.1+/-160.7 | | | 133.1+/-160.7 | |
| _ | | | 339.26+/-339.95 | | | 353.59+/-366.06 | |
| | | | N1-40, N2=168 | × | × | N1=40, N2=31 | × |
| _ | | | Fold Change: 2.57 | | | Fold Change: 2.7 | |
| | | | P-value: .00002 | | | P-value: .00061 | |
| 196 | 393 | AI218358 | 210.74+/-117.83 | | 210.74+/-117.83 | 210.74+/-117.83 | 210.74+/-117.83 |
| | | | 87.3+/-105.07 | | 82.51+/-91.18 | 76.3+/-97.56 | 74.44+/-133.85 |
| | | | N1=39, N2=168 | × | N1=39, N2=10 | N1=39, N2=31 | N1=39, N2=6 |
| _ | | | Fold Change: 2.43 | | Fold Change: 2.74 | Fold Change: 2.64 | Fold Change: 3.62 |
| | | | P-value: 0 | | P-value: .0333 | P-value: 0 | P-value: .00101 |

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|-----|--------|----------|-------------------|--|---------------------------|--------------------|---------------------|
| * | Sed ID | Genbank | Normal vs All | Normal vs Malignant | Normal vs Stage I | Normal vs Stage II | Normal vs Stage III |
| 197 | 394 | AX222594 | 431.73+/-162.38 | | | 431.73+/-162.38 | |
| | | | 200 001 110 40 | | | 140 711 01 02 | |
| | | | 109.82+/-117.45 | | | 143.747-81.33 | |
| | | | N1=40, N2=168 | × | × | N1=40, N2=31 | × |
| | | | Fold Change: 2.81 | | | Fold Change: 3.23 | |
| | | | P-value: 0 | | | P-value: 0 | |
| 861 | 397 | AI242160 | 320,5+/-159,05 | | | | |
| | | | 170.68+/-74.58 | | | | |
| | | | N1=30 N2=168 | ۲ | Þ | × | × |
| | | | Pold Change: 1 72 | • | 4 | | 4 |
| | | | P-value: 0 | | | | |
| 8 | 398 | AT244908 | | Yes a second sec | | | 425.47+/-356.02 |
| | | | | | | | 148 1+/-83 25 |
| | | | Þ | Þ | Þ | > | M1=40 N2=10 |
| | | | < | < | 4 | 4 | Told Change: 2 51 |
| | | | | | | | P-value: 00165 |
| 900 | 300 | AT247827 | 250 231/ 21/ 52 | | | 250 33+1,314 52 | 250 33+/-314 52 |
| 3 | | | 40.01+/-55.99 | | | 34 79+/-48 59 | 20.13+/-36.3 |
| | | | N1=40 N2=169 | × | > | N1=40 N2=31 | N1=40 N2=10 |
| | | | Told Change: 2 99 | ¢ | 4 | Fold Change: 4 12 | Fold Change: 4 40 |
| | | | P-volue: 0 | | | Parahar 0 | D_value: 0 |
| 1 | | | T dino. | | | - Animo | o commit T |
| 707 | 401 | AI264135 | | | | | 246.38+/-130.33 |
| | | | | | | | 97.95+/-69.13 |
| | | | × | × | × | × | N1=40, N2=10 |
| | | | | | | | Fold Change: 2.68 |
| | | | | | | | P-value: .00126 |
| 202 | 403 | AI266650 | | | | | 657.85+/-222.05 |
| | | | | | | | 309.8+/-83.6 |
| | | | × | × | × | × | N1=40, N2=10 |
| | | | | | | | Fold Change: 2.09 |
| | | | | | | | P-value: ,00001 |
| 203 | 405 | AI275140 | | | | 26.4+/-47.96 | |
| | | | | | | 232.81+/-210.09 | |
| | | | × | × | × | N1=40, N2=31 | × |
| | | | | | | Fold Change: 641 | |
| | | | | | | P-value: 0 | |

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|-----|--------|----------|-------------------|---------------------|-------------------|--------------------|---------------------|
| # | Seq ID | Genbank | Normal vs All | Normal vs Malignant | Normal vs Stage I | Normal vs Stage II | Normal vs Stage III |
| 204 | 407 | A1276259 | 121.2+/-221.34 | | | 121.51+/-224.22 | |
| | | | 238.08+/-274.65 | | | 245.37+/-268.39 | |
| | | | N1=40, N2=168 | × | × | N1=40, N2=31 | × |
| | | | Fold Change: 2.02 | | | Fold Change: 2.09 | |
| | | | P-value: .00019 | | | P-value: .00466 | |
| 205 | 408 | AI277612 | 1009,49+/-899,38 | | 1022.91+/-907.07 | 1022.91+/-907.07 | 1022.91+/-907.07 |
| | ! | | 93.86+/-163.03 | | 163.73+/-156.36 | 82.75+/-117.04 | 40.07+/-27.26 |
| | | | N1=40, N2=168 | × | N1=40, N2=6 | N1=40, N2=31 | N1=40, N2=10 |
| | | | Fold Change: 9.23 | | Fold Change: 5.12 | Fold Change: 10.43 | Fold Change: 16.08 |
| | | | P-value: 0 | | P-value: .00592 | P-value: 0 | P-value: 0 |
| 206 | 413 | AI285970 | | | | 269.01+/-274.71 | |
| | | | | | | 88.12+/-54.51 | |
| | | | × | × | × | N1=40, N2=31 | × |
| | | | | | | Fold Change: 2 | |
| | | | | | | P-value: .00393 | |
| 207 | 414 | A7288586 | | | | 313,32+/-139,2 | 313.32+/-139.2 |
| | | | | | | 113.394/-56.77 | 107.36+/-49.96 |
| | | | × | × | × | N1=39, N2=31 | N1=39, N2=6 |
| | | | | | | Fold Change: 2.6 | Fold Change: 2.94 |
| | | | | | | P-value: 0 | P-value: .00028 |
| 208 | 415 | AI288745 | 276.49+/-146.02 | | | 276.49+/-146.02 | 276.49+/-146.02 |
| _ | | | 108.9+/-73.82 | | | 104.75+/-63.19 | 67.42+/-24.06 |
| | | | N1=39, N2=168 | × | × | N1=39, N2=31 | N1=39, N2=6 |
| | | | Fold Change: 2.51 | | - | Fold Change: 2.5 | Fold Change: 3.69 |
| | | | P-value: 0 | | | P-value: 0 | P-value: 0 |
| 209 | 417 | AI300876 | 601.83+/-985.51 | | | 601.83+/-985.51 | 601.83+/-985.51 |
| | | | 74.02+/-258.41 | | | 54.33+/-149.24 | 13.6+/-22.13 |
| | | | N1=40, N2=168 | × | × | N1=40, N2=31 | N1=40, N2=10 |
| | | | Fold Change: 6.47 | | | Fold Change: 6.66 | Fold Change: 8.67 |
| | | | P-value: 0 | | | P-value: 0 | P-value: 0 |
| 210 | 422 | AI333767 | | | | | 201.68+/-104.32 |
| | | | | | | | 104.93+/-75.04 |
| | | | × | × | × | × | N1=40, N2=10 |
| | | | | | | | Fold Change: 2 |
| | | | | , | | | P-value: .00805 |

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|-----|--------|----------|--|---------------------|--|--|--|
| # | Seq ID | Genbank | Normal vs All | Normal vs Malignant | Normal vs Stage I | Normal vs Stage II | Normal vs Stage III |
| 211 | 423 | AI333987 | × | × | × | 208.53+/-320.79 57.06+/-100.56 N1=40, N2=31 Fold Change: 2.4 P-value: .00125 | × |
| 212 | 424 | AI338536 | × | × | × | × | 548.6+/-192.05 227.52+/-68.12 N1=40, N2=10 Fold Change: 2.31 P-value: 0 |
| 213 | 427 | AI341602 | 135.164/-276.86 593.584/-944.91 N1=40, N2=168 Fold Change: 4.46 P-value: 0 | × | 137.44+/-280.1 438.85+/-447.12 N1=40, N2=6 Fold Change: 3.86 P-value: .03152 | 137.44+/-280.1 723.06+/-652.3 N1==40, N2==31 Fold Change: 6.28 P-value: 0 | 137.44+/-280.1 580.91+/-380.87 N1=40, N2=10 Fold Change: 6.09 P-value: .00062 |
| 214 | 428 | AI342169 | × | × | × | × | 496.19+/-393.99 1347.53+/-986.09 N1=40, N2=10 Fold Change: 2.64 P-value: .0043 |
| 215 | 430 | AI344312 | 84.72+/-57.63 209.71+/-134.29 N1=40, N2=168 Fold Change: 2.33 P-value: 0 | × | × | × | × |
| 216 | 431 | AI346341 | 640.59+/-422.41 155.96+/-203.21 N1=40, N2=168 Fold Change: 4.26 P-value: 0 | × | × | 635.18+/-426.52 120.31+/-143.98 N1=40, N2=31 Fold Change: 4.92 P-value: 0 | 635.18+/-426.52 104.05+/-133.87 N1=40, N2=10 Fold Change: 5.89 P-value: .00006 |
| 217 | 432 | AI351043 | 236.73+/-93.44 109.27+/-56.02 N1=40, N2=168 Fold Change: 2.23 P-value: 0 | × | × | 236.3+/-94.62 98.05+/-50.78 N1=40, N2=31 Fold Change: 2.47 P-value: 0 | 236.3+/-94.62 73.16+/-38.18 N1=40, N2=10 Fold Change: 3.35 P-value: .00009 |

| H az | .28 | 2 Y | 2.86 | 21 | 22.1 | 10 | 2.51 | | .11 | 28 | φ | 1.82 | (18 | :58 | 80 | 10 | 3.09 | _ | .18 | 17 | 10 | 2.08 | _ | 66'(| 1.5 | 10 | 10 | 2 | 2 | 96. | 2 5 | 2 |
|---------------------|-----------------|-------------------------------|-------------------|----------------|----------------|--------------|-------------------|------------|----------------|----------------|--------------|-------------------|-----------------|-----------------|----------------|---------------|-------------------|------------|----------------|----------------|---------------|-------------------|------------|-----------------|----------------|--------------|--------------|-------------------|-----------------|-----------------|--------------------------------|-------|
| Normal vs Stage III | 222.69+/-144.28 | 51.96+/-71.13 N1=39. N2=6 | Fold Change: 2.86 | 103.44+/-82.21 | 229.26+/-76.25 | N1=40, N2=10 | Fold Change: 2.51 | P-value: 0 | 209.31+/-66.11 | 116.1+/-36.28 | N1=39, N2=6 | Fold Change: 1.82 | P-value: .00118 | 404.52+/-178.58 | 124.99+/48.08 | N1=40, N2=10 | Fold Change: 3.09 | P-value: 0 | 211.26+/-81.18 | 95.25+/-21.17 | N1=40, N2=10 | Fold Change: 2.08 | P-value: 0 | 360.06+/-360.99 | 160.98+/-94.5 | N1=40, N2=10 | N1=40, N2= | Fold Change: 2 | P-value: .0091 | 815.22+/-371.96 | N1=40 N2=10 | A |
| Normal vs Stage II | 222.69+/-144.28 | 77.49+/-85.14 N1=39. N2=31 | Fold Change: 2.28 | 7-4 | | × | | | 226.33+/-66.64 | 131.77+/-56.91 | N1=39, N2=31 | Fold Change: 1.8 | P-value: 0 | 404.52+/-178.58 | 156.69+/-71.33 | N1=40, N2=31 | Fold Change: 2.52 | P-value: 0 | 211.26+/-81.18 | 101.07+/-42.95 | N1=40, N2=31 | Fold Change: 2.07 | P-value: 0 | 360,06+/-360,99 | 150.11+/-76.77 | N1=40, N2=31 | N1=40, N2=31 | Fold Change: 2.07 | P-value: .00001 | 815.22+/-371.96 | N1=40 N2=31 | ***** |
| Normal vs Stage I | 0 | × | | | | × | | | | | × | | | | | × | | | | | × | | | | | × | × | | | | > | • |
| Normal vs Malienant | | × | | | | × | | | | | × | | | | | × | | | | | × | | | | | × | × | | | | * | • |
| Normal vs All | | * | l | | | × | | | | | × | | | 412.08+/-182.64 | 160.8+/-88.56 | N1=40, N2=168 | Fold Change: 2.56 | P-value: 0 | 215.71+/-84.93 | 107.21+/-43.49 | N1=40, N2=168 | Fold Change: 2.02 | P-value: 0 | | | × | × | | | 837.48+/-393.22 | 288.6+/-150.82 N1=40 N2=168 | 100 |
| Genbank | AI352171 | | | AT347630 | Con located | | | | AI361002 | | | | | AI369275 | | | | | AI375115 | | | | | AX377937 | | | | | | AI378584 | | |
| Sea ID | 434 | | | 436 | 2 | | | | 437 | | | | | 14 | | | | | 443 | | | | | 446 | | | | | | 447 | | |
| * | 218 | | | 210 | ì | | | | 220 | | | | | 221 | | | | | 222 | | | | | 223 | | _ | | | | 224 | | |

| г | _ | | | _ | | _ | _ | _ | _ | | | - | _ | _ | _ | | _ | | | _ | | _ | | _ | | _ | _ | | | - | _ | | _ | _ | _ |
|---|---------------------|-----------------|-----------------|---------------|-------------------|-----------------|-----------------|----------------|--------------|-------------------|-----------------|----------------|-----------------|--------------|-------------------|-----------------|-----------------|----------------|---------------|-------------------|-----------------|---------------|-----------------|--------------|-------------------|----------------|-----------------|--------------|-------------------|-----------------|-----------------|-----------------|---------------|-------------------|-----------------|
| | Normal vs Stage III | 380.22+/-173.64 | 68.16+/-48.01 | N1=40, N2=10 | Fold Change: 5.51 | P-value: 0 | | | × | | | 191,48+/-96.27 | 387.31+/-138.62 | N1=40, N2=10 | Fold Change: 2.16 | P-value: .00071 | 269.91+/-245.06 | 62.78+/-171.44 | N1=39, N2=6 | Fold Change: 3.91 | P-value: .0021 | | | × | | | | × | | | 933.35+/-487.41 | 240.55+/-123.45 | N1=40, N2=10 | Fold Change: 3.74 | P-value: .00001 |
| | Normal vs Stage II | 380.22+/-173.64 | 87.51+/-60.3 | N1-40, N2=31 | Fold Change: 4.24 | P-value: 0 | 194.43+/-169.86 | 361,5+/-246,51 | N1=40, N2=31 | Fold Change: 2.15 | P-value: .00078 | | | × | | | 269.91+/-245.06 | 34.22+/-46.82 | NI=39, N2=31 | Fold Change: 3.85 | P-value: 0 | 74.44+/-64.03 | 339.43+/-658.29 | N1=40, N2=31 | Fold Change: 2.18 | 72 16±/.151 56 | 408 66+/-668 16 | N1=40, N2=31 | Fold Change: 2.63 | P-value: .00745 | 933.35+/-487.41 | 238.15+/-85.44 | N1=40, N2=31 | Fold Change: 3.54 | P-value: 0 |
| | Normal vs Stage I | 380.22+/-173.64 | 136.6+/-134.47 | N1-40, N2-6 | Fold Change: 3.5 | P-value: .03454 | | | × | | | | | × | | | 269.91+/-245.06 | 63.86+/-29.55 | N1=39, N2=10 | Fold Change: 3.2 | P-value: .00133 | | | × | | | | × | | | 933.35+/-487.41 | 387.97+/-131.13 | N1=40, N2=6 | Fold Change: 2.11 | P-value: .0003 |
| | Normal vs Malignant | | | × | | | | | × | | | | | × | | | | | × | | | | | × | | | | × | | | | | × | | |
| | Normal vs All | 386.86+/-176.47 | 126,18+/-143.63 | N1=40, N2=168 | Fold Change: 3.46 | P-value: 0 | | | × | | | | | × | | | 269,91+/-245.06 | 44,47+/-65,54 | N1=39, N2=168 | Fold Change; 3.7 | P-value: 0 | | | × | | | | × | | | 927.3+/-482.64 | 259.61+/-119.48 | N1=40, N2=168 | Fold Change: 3.36 | P-value: 0 |
| | Genbank | AI379723 | | | | | AI380204 | | | | | AI380932 | | | | | AT381930 | | | | | AI393356 | | | | AT401024 | 7C01047W | | | | AI417267 | | | | |
| | Sed ID | 448 | | | | | 450 | | | | | 452 | | | | | 456 | | | | | 458 | | | | 127 | 401 | | | | 462 | | | | |
| | # | 225 | | | | | 226 | | | | | 227 | <u> </u> | | | | 228 | | | | | 229 | | | | 35 | 67 | | _ | | 231 | | | | |

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|-----|--------|----------|-------------------|---------------------|-------------------|---------------------------------|---------------------|
| | Seq ID | Genbank | Normal vs All | Normal vs Malignant | Normal vs Stage I | Normal vs Stage II | Normal vs Stage III |
| 232 | 463 | AI417917 | 564.39+/-315.14 | | | 567.47+/-318.65 | 567.47+/-318.65 |
| | | | 271 17+/-167 23 | | | 218 33+/-110.93 | 115.26+/-82.27 |
| | | | N1=40, N2=168 | × | × | N1-40, N2=31 | N1=40. N2=10 |
| | | | Fold Change: 2 25 | | | Fold Change: 2.61 | Fold Change: 5.16 |
| | | | P-value: 0 | | | P-value: 0 | P-value: .00005 |
| 233 | 465 | AI418481 | | | | 77.52+/-60.1 | |
| | | | | | | 257.35+/-253.01 | |
| | | | × | × | × | N1=40, N2=31 | × |
| | | | | | | Fold Change: 2.83 P-value: 0 | |
| 234 | 466 | AI418596 | | | | | 229.67+/-120.12 |
| | | | | | | | 43.69+/-43.08 |
| | | | × | × | × | × | N1=39. N2=6 |
| | | | | | | | Fold Change: 3.79 |
| | | | | | | | P-value: 0 |
| 235 | 467 | AI419030 | 446,12+/-255.78 | | | 445.97+/-259.12 | 445.97+/-259.12 |
| | | | 158.41+/-132.36 | | | 133.67+/-123.4 | 81.99+/-31.49 |
| | | | N1=40, N2=168 | × | × | N1=40, N2=31 | N1=40, N2=10 |
| | | | Fold Change: 3.03 | | | Fold Change: 3.56 | Fold Change: 4.69 |
| | | | P-value: 0 | | | P-value: 0 | P-value: 0 |
| 336 | 468 | A1421837 | 294.7+/-145.9 | | | 293.96+/-147.73 | 293.96+/-147.73 |
| | | | 111.04+/-64.19 | | | 102.42+/-57.12 | 109.78+/-55.53 |
| | | | N1=40, N2=168 | × | × | N1=40, N2=31 | N1=40, N2=10 |
| | | | Fold Change: 2.62 | | | Fold Change: 2.8 | Fold Change: 2.62 |
| | | | P-value: 0 | | | P-value: 0 | P-value: .00166 |
| 237 | 469 | A1431799 | | | 463.05+/-382.43 | | |
| | | | | | 658.8+/-240.03 | | |
| | | | × | × | N1=40, N2=6 | × | × |
| | | | | | Fold Change: 2 | | |
| | | | | | P-value: .00993 | | |
| 238 | 471 | AI435828 | | | | 587.69+/-414.14 | 587.69+/-414.14 |
| | | | | | | 299.14+/-333.14 | 119.57+/-42.81 |
| | | | × | × | × | N1=40, N2=31 | N1=40, N2=10 |
| | | | | | | Fold Change: 2.52 | Fold Change: 3.9 |
| | | | | | | P-value: .00061 | P-value: .00001 |

| * | Seq ID | Genbank | Normal vs All | Normal vs Malignant | Normal vs Stage I | Normal vs Stage II | Normal vs Stage III |
|-----|--------|----------|-------------------|---------------------|-------------------|--------------------|---------------------|
| 239 | 475 | AI446030 | 383.04+/-264.2 | | | 383.04+/-264.2 | |
| | | | 025 52±7.016 21 | | | 987 66±1,-795 06 | |
| | | | N1=40 N2=168 | * | × | N1=40 N2=31 | × |
| | | | Fold Change: 2.29 | 1 | | Fold Change: 2.53 | |
| | | | P-value: 0 | | | P-value: 0 | |
| 240 | 476 | AI446168 | 216.67+/-584.58 | | | 220,39+/-591,75 | 220,39+/-591.75 |
| | | | 561.08+/-695.63 | | | 606,41+/-908,81 | 603.11+/-527.41 |
| | | | N1=40, N2=168 | × | × | N1=40, N2=31 | N1=40, N2=10 |
| | | | Fold Change: 3.28 | | | Fold Change: 3.51 | Fold Change: 3.91 |
| _ | | | P-value: 0 | | | P-value: 0 | P-value: .00476 |
| 241 | 477 | AI458003 | 276.51+/-201.47 | | 280.16+/-202.76 | 280.16+/-202.76 | 280.16+/-202.76 |
| | | | 38.41+/-97.2 | | 81.96+/-89.4 | 43.42+/-152.26 | 4.43+/-42.79 |
| | | | N1=40, N2=168 | × | N1=40, N2=6 | N1=40, N2=31 | N1=40, N2=10 |
| | | | Fold Change: 5.07 | | Fold Change: 3.22 | Fold Change: 4.35 | Fold Change: 7.36 |
| | | | P-value: 0 | | P-value: .03405 | P-value: 0 | P-value: 0 |
| 242 | 482 | A1468491 | | | | | 256.57+/-264.41 |
| | | | | | | | 84.09+/-113.95 |
| | | | × | × | × | × | N1=40, N2=10 |
| | | | | | | | Fold Change: 2.68 |
| | | | | | | | P-value: .00796 |
| 243 | 487 | AI480215 | 122.23+/-104.86 | | | 125.66+/-103.93 | |
| | | | 295.22+/-297.68 | | | 321.02+/-262.5 | |
| | | | N1=40, N2=168 | × | × | N1=40, N2=31 | × |
| | | | Fold Change: 2.15 | | | Fold Change: 2.49 | |
| | | | P-value: .00005 | | | P-vame: .0002 | |
| 244 | 489 | AI492051 | 384,45+/-175.99 | | | 382.34+/-177.78 | 382.34+/-177.78 |
| | | | 157,89+/-176.84 | | | 91.96+/-44.92 | 154.08+/-202.28 |
| | | | N1=40, N2=168 | × | × | N1=40, N2=31 | N1-40, N2-10 |
| | | | Fold Change: 3,12 | | | Fold Change: 4.08 | Fold Change: 3.91 |
| | | | P-value: 0 | | | P-value: 0 | P-value: .0032 |
| 245 | 490 | AI492091 | | | | | 146.39+/-59.05 |
| | | | | | | | 321.61+/-116.38 |
| | | | × | × | × | × | N1=40, N2=10 |
| | | | | | | | Fold Change: 2.22 |
| _ | | | | | | | P-value: .00005 |

| 4 | SeqID | Genbank | Normal vs All | Normal vs Malignant | Normal vs Stage I | Normal vs Stage II | Normal vs Stage III |
|-----|-------|----------|-------------------|---------------------|-------------------|--------------------|---------------------|
| 246 | 493 | AI492879 | 214.23+/-650.46 | | 219.42+/-658.12 | 219.42+/-658.12 | 219.42+/-658.12 |
| | | | 465.24+/-396.65 | | 277.88+/-265.92 | 663.74+/-488.43 | 771.71+/-307.65 |
| | | | N1=40, N2=168 | × | N1=40, N2=6 | N1=40, N2=31 | N1=40, N2=10 |
| | | | Fold Change: 6.22 | | Fold Change: 3.72 | Fold Change: 9.89 | Fold Change: 14.12 |
| | | | P-value: 0 | | P-value: .01903 | P-value: 0 | P-value: 0 |
| 247 | 494 | AI498375 | 244,41+/-203,11 | | 244.41+/-203.11 | 244.41+/-203.11 | 244.41+/-203.11 |
| | | | 46+/-55.84 | | 47.6+/-45.95 | 43.16+/-57.68 | 48.55+/-41.9 |
| | | | N1=39, N2=168 | × | N1=39, N2=10 | N1=39, N2=31 | N1=39, N2=6 |
| | | | Fold Change: 3.62 | | Fold Change: 3.32 | Fold Change: 3.62 | Fold Change: 3.45 |
| | | | P-value: 0 | | P-value: .00134 | P-value: 0 | P-value: .00002 |
| 248 | 498 | AI499334 | 236.34+/-181.29 | | 236.34+/-181.29 | | 236,34+/-181,29 |
| | | | 117.03+/-221.42 | | 83.26+/-88.81 | | 16.68+/-51.22 |
| | | | N1=39, N2=168 | × | N1=39, N2=10 | × | N1=39, N2=6 |
| | | | Fold Change: 2.28 | | Fold Change: 2.68 | | Fold Change: 4.22 |
| | | | P-value: 0 | | P-value: .0244 | | P-value: 0 |
| 249 | 200 | AI524085 | 380.84+/-525.16 | | 388.89+/-529.52 | 388.89+/-529.52 | 388.89+/-529.52 |
| | | | 54.03+/-109.2 | | 35.06+/-63.65 | 36.84+/-79.16 | 8.33+/-48.11 |
| | | | N1=40, N2=168 | × | N1=40, N2=6 | N1=40, N2=31 | N1=40, N2=10 |
| | | | Fold Change: 4.48 | | Fold Change: 4.76 | Fold Change: 5.39 | Fold Change: 7 |
| | | | P-value: 0 | | P-value: .00065 | P-value: 0 | P-value: 0 |
| 250 | 501 | AIS25044 | 319,4+/-142.13 | | | 316.89+/-143.08 | 316.89+/-143.08 |
| | | | 131.06+/-178.35 | | | 121.55+/-76.85 | 65.73+/-20.84 |
| | | | N1=40, N2=168 | × | × | N1=40, N2=31 | N1=40, N2=10 |
| | | | Fold Change: 2.93 | | | Fold Change: 2.83 | Fold Change: 4.61 |
| | | | P-value: 0 | | | P-value: 0 | P-value: 0 |
| 251 | 205 | AI525601 | | | | | 223.25+/-101.98 |
| | | | | | | | 605.18+/-406.21 |
| | | | × | × | × | × | N1=40, N2=10 |
| | | | | | | | Fold Change: 2.43 |
| | | | | | | | P-value: .00221 |
| 252 | 504 | AI535997 | 526.38+/-278.8 | | | 527.18+/-282.39 | 527.18+/-282.39 |
| | | | 203,43+/-127.91 | | | 182.64+/-99.43 | 177.05+/-109.73 |
| | | | N1=40, N2=168 | × | × | N1=40, N2=31 | N1=40, N2=10 |
| | | | Fold Change: 2.55 | | | Fold Change: 2.76 | Fold Change: 2.81 |
| | | | P-value: 0 | | | P-vaine: 0 | P-value: .00005 |

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|-----|--------|----------|--------------------|---------------------|--|--------------------|----------------------|
| | Seq ID | Genbank | Normal vs All | Normal vs Malignant | Normal vs Stage I | Normal vs Stage II | Normal vs Stage III |
| 253 | 206 | AI539386 | 1911.09+/-2400.57 | | | 1924.9+/-2430.34 | |
| | | | 4189 96+/-4852 35 | | | 4667 18+/-4736 01 | |
| | | | 40.00 | ; | ; | 10 010 01 | ; |
| | | | NI=40, NZ=168 | * | * | NI=40, NZ=51 | * |
| | | | Fold Change: 2.2 | | | Fold Change: 2.62 | |
| | | | P-value: .00002 | | | P-value: .00005 | |
| 254 | 507 | AI539443 | 105.57+/-75.36 | | | 105.57+/-75,36 | |
| | | | 244 51+/-212 14 | | | 235 02+1-188 35 | |
| | | | NT-20 ND-160 | Þ | Þ | NI1-20 NO-21 | ٥ |
| | | | 141 -00, 142 -100 | 4 | 4 | 10 24 60 14 | 4 |
| | | | Fold Change: 2.05 | | | Fold Change: 2.11 | |
| | | | P-value: 0 | | | P-value: .00011 | |
| 255 | 509 | AI546943 | | | | | 310.79+/-204.98 |
| | | | | | | | 120 98+/-130 81 |
| | | | Þ | Þ | Þ | Þ | N1-40 ND-10 |
| | | | * | * | * | < | NI=70 N7=10 |
| | | | | | | | Fold Change: 3.17 |
| | | | | | | | P-value: .01143 |
| 256 | 510 | AI553918 | 831.82+/-279.85 | | | 821.96+/-276.38 | 821.96+/-276.38 |
| | | | 384.25+/-176.84 | | | 364.75+/-169.81 | 378.99+/-216.53 |
| | | | N1=40, N2=168 | × | × | N1=40, N2=31 | N1=40. N2=10 |
| | | | Told Changes: 2 22 | | | Fold Change: 7 21 | Told Change: 2 42 |
| | | | Fold Change, 2.22 | | | Total Change: 2:01 | Total Culture Contra |
| - 1 | | | F-value: 0 | | | r-vaine: 0 | F-value: .00515 |
| 257 | 511 | AI554514 | | | 90.74+/-52.8 232.59+/-162.79 | | |
| | | | × | × | N1=40. N2=6 | × | × |
| | | | | | Fold Change: 2.54 | | |
| 258 | 512 | AI557210 | 126.32+/-140.31 | | | 129.15+/-140.98 | 129.15+/-140.98 |
| | | | 534.58+/-397.4 | | | 364.46+/-223.6 | 306.19+/-192.07 |
| | | | N1=40. N2=168 | × | × | N1=40. N2=31 | N1=40. N2=10 |
| | | | Fold Change: 4 09 | | | Fold Change: 3 55 | Fold Change: 2 84 |
| | | | D malana | | | D meline. | D 110 110 00015 |
| 1. | П | · Second | r-value. o | | | O Ammo. O | C1 C0C : 00 200 |
| 60 | cic | ALSOUND4 | | | | 683.27/-300.13 | 663.24/-200.13 |
| | | | | | | 473,3+/-341.89 | 284.56+/-117.13 |
| | | | × | × | × | N1-40, N2-31 | N1=40, N2=10 |
| | | | | | | Fold Change: 2.17 | Fold Change: 3.2 |
| | | | | | | P-value: 0 | P-value: .00002 |

| * | Seq ID | Genbank | Normal vs All | Normal vs Malignant | Normal vs Stage I | Normal vs Stage II | Normal vs Stage III |
|-----|--------|----------|-------------------|---------------------|-------------------|--------------------|---------------------|
| 260 | 919 | AIS60159 | | | | 249.15+/-159.54 | 249.15+/-159.54 |
| | | | × | × | × | N1=40, N2=31 | N1=40, N2=10 |
| | | | 1 | | | Fold Change: 2.08 | Fold Change: 3.13 |
| | | | | | | P-value: 0 | P-value: .00061 |
| 261 | 517 | A1566038 | 257.62+/-109.32 | | | 436.1+/-150.59 | |
| | | | 154.67+/-104.12 | | | 302.74+/-153.34 | |
| | | | N1=39, N2=168 | × | × | N1=39, N2=31 | × |
| | | | Fold Change: 1.94 | | | Fold Change: 1.57 | |
| | | | P-value: 0 | | | P-value: .00104 | |
| 797 | 523 | AI583942 | 231.62+/-1223.34 | | | 237,554/-1238.75 | |
| | | | 574.26+/-2305.76 | | | 1788.59+/-4779.29 | |
| | | | N1=40, N2=168 | × | × | N1=40, N2=31 | × |
| | | | Fold Change: 2.18 | | | Fold Change: 4.69 | |
| | | | P-value: .00076 | | | P-value: .00099 | - |
| 263 | 524 | AT587178 | | | | | 222.78+/-143.35 |
| 3 | | | | | | | 88.43+/-75.43 |
| | | | × | × | × | × | N1=40, N2=10 |
| | | | | | | | Fold Change: 2.64 |
| | | | | | | | P-vaine: .00579 |
| 264 | 526 | A1589858 | 296.87+/-162.49 | | | 296.87+/-162.49 | 296.87+/-162.49 |
| | | | 138.954/-123 | | | 143.85+/-108.99 | 121.61+/-47.86 |
| | | | N1=39, N2=168 | × | × | N1-39, N2-31 | N1=39, N2=6 |
| | | | Fold Change: 2.17 | | | Fold Change: 2.1 | Fold Change: 2.31 |
| | | | P-value: 0 | | | P-value: .00002 | P-value: .00132 |
| 265 | 527 | AI590093 | 316.95+/-171.55 | | | 314.33+/-172.98 | 314.33+/-172.98 |
| | | | 167.02+/-141.99 | | | 149,67+/-84.85 | 121.07+/-88.54 |
| | | | N1=40, N2=168 | × | × | N1=40, N2=31 | N1=40, N2=10 |
| | | | Fold Change: 2.1 | | | Fold Change: 2.08 | Fold Change: 2.91 |
| | | | P-value: 0 | | | P-value: .00003 | P-value: .00504 |
| 566 | 529 | AI598252 | | | | 417.54+/-196.25 | |
| | | | ; | ; | , | 833.81+/-317.10 | Þ |
| | | | × | × | * | NI=59, NZ=51 | < |
| | | | | | | Fold Change: 2.00 | |
| | | | | | | L-Value, o | |

| * | Seq ID | Genbank | Normal vs All | Normal vs Malignant | Normal vs Stage I | Normal vs Stage II | Normal vs Stage III |
|-----|--------|----------|-------------------|---------------------|-------------------|--------------------|--|
| 267 | 530 | AI601149 | 270.5+/-122.37 | | | 267.15+/-122.1 | 267.15+/-122.1 |
| | | | 115.84+/-107.17 | | | 101,95+/-53.86 | 55.7+/-20.35 |
| | | | N1=40, N2=168 | × | × | N1=40, N2=31 | N1=40, N2=10 |
| | | | Fold Change: 2.62 | | | Fold Change: 2.69 | Fold Change: 4.57 |
| | | | P-value: 0 | | | P-value: 0 | P-value: 0 |
| 268 | 532 | AI610837 | 2072.69+/-1692.17 | | | 2072.69+/-1692.17 | 2072.69+/-1692.17 |
| | | | 724.59+/-659.17 | | | 544+/-358.27 | 319.79+/-159.71 |
| | | | N1=40, N2=168 | × | × | N1=40, N2=31 | N1=40, N2=10 |
| | | | Fold Change: 2.65 | | | Fold Change: 3.17 | Fold Change: 4.66 |
| | | | P-value: 0 | | | P-value: 0 | P-value: 0 |
| 569 | 534 | AI620381 | | | | | 610.77+/-316.3 |
| | | | | | | | 1217.36+/-274.04 |
| | | | × | × | × | × | N1=40, N2=10 |
| | | | | | | | Fold Change: 2.16 |
| | | | | | | | P-value: 0 |
| 270 | 535 | AI624103 | 289.59+/-159.89 | | 289.59+/-159.89 | 289.59+/-159.89 | 289.59+/-159.89 |
| | | | 123.14+/-143.73 | | 116.4+/-81.91 | 82.42+/-59.19 | 71.19+/-34.34 |
| | | | N1=39, N2=168 | × | N1=39, N2=10 | N1=39, N2=31 | N1=39, N2=6 |
| | | | Fold Change: 2.74 | | Fold Change: 2.65 | Fold Change: 3.4 | Fold Change: 3.74 |
| | | | P-value: 0 | | P-value: .03426 | P-value: 0 | P-value: 0 |
| 271 | 537 | AI631301 | 328.42+/-165.41 | | | 328.42+/-165.41 | 328.42+/-165.41 |
| | | | 126.68+/-74.53 | | | 111+/-70.99 | 58.7+/-32.01 |
| | | | N1=39, N2=168 | × | × | N1=39, N2=31 | N1=39, N2=6 |
| | | | Fold Change: 2.52 | | | Fold Change: 2.8 | Fold Change: 5.02 |
| | | | P-value: 0 | | | P-value: 0 | P-value: 0 |
| 272 | 538 | AI631850 | | | | 26.13+/-35.18 | |
| | | | | | | 224.44+/-595.61 | |
| | | | × | × | × | N1=40, N2=31 | × |
| | | | | | | Fold Change: 2.14 | |
| | | | | | | P-value: .00784 | The second secon |
| 273 | 240 | AI634852 | | | | 278.07+/-162.92 | |
| | | | × | × | × | N1=40, N2=31 | × |
| | | | | | | Fold Change: 2.18 | |
| | | | | | | P-value: .0011 | |

| 7 | Good III | Control | Normal ve All | Normal ve Malianant | Normal ve Stage | Normal ve Stage II | Normal ve Stage III |
|-----|----------|----------|-------------------|------------------------|-------------------|--------------------|---------------------|
| * | or hac | Gembaur | LIGHT NA PAIL | TOT HERE AS TARRESTAND | TOTAL 13 DIRECT | Troumer to Singery | TOTAL IN COURSE |
| 274 | 541 | AI635774 | 212.33+/-93.64 | | | 212,33+/-93.64 | 212.33+/-93.64 |
| | | | 92.03+/-51.47 | | | 93.14+/-61.28 | 89.05+/-56.69 |
| | | | N1=39, N2=168 | × | × | N1=39, N2=31 | N1=39, N2=6 |
| | | | Fold Change: 2.39 | | | Fold Change: 2.5 | Fold Change: 2.62 |
| | | | P-value: 0 | | | P-value: 0 | P-value: .00169 |
| 275 | 546 | AI650514 | 108,33+/-162,01 | | 110.57+/-163.5 | 110.57+/-163.5 | 110.57+/-163.5 |
| | | | 321.96+/-278.48 | | 306.38+/-174.2 | 495.16+/-349.41 | 380.44+/-266.81 |
| | | | N1=40, N2=168 | × | N1=40, N2=6 | N1=40, N2=31 | N1=40, N2=10 |
| | | | Fold Chance: 3.14 | | Fold Change: 3.3 | Fold Change: 5.31 | Fold Change: 3,44 |
| | | | P-value: 0 | | P-value: .02515 | P-value: 0 | P-value: .01035 |
| 276 | 550 | AI651732 | | | | | 200.01+/-105.06 |
| | | | | | | | 67.38+/-39.88 |
| | | | × | × | × | × | N1=40, N2=10 |
| | | | | | | | Fold Change: 3.01 |
| | | | | | | | P-value: .00028 |
| 277 | 551 | AI652058 | | | | | 182.73+/-51.39 |
| | | | | | | | 384.98+/-89.44 |
| | | | × | × | × | × | N1=40, N2=10 |
| | | | | | | | Fold Change: 2.14 |
| | | | | | | | P-value: 0 |
| 278 | 552 | AI652459 | | | | | 724.8+/-344.86 |
| | | | | | | | 274.16+/-175.44 |
| | | | × | × | × | × | N1=40, N2=10 |
| | | | | | | | Fold Change: 2.8 |
| | | | | | | | P-value: .00159 |
| 279 | 554 | AI653487 | 251.37+/-223.37 | | 251.37+/-223.37 | 251.37+/-223.37 | 251.37+/-223.37 |
| | | | 44.78+/-81.85 | | 41.71+/-73.61 | 30.69+/-76.04 | 4.44+/-69.04 |
| | | | N1=39, N2=168 | × | N1=39, N2=10 | N1=39, N2=31 | N1=39, N2=6 |
| | | | Fold Change: 3.04 | | Fold Change: 3.36 | Fold Change: 3.36 | Fold Change: 4.16 |
| | | | P-value: 0 | | P-value: .00441 | P-value: 0 | P-value: .00002 |
| 280 | 555 | AI654035 | 227.06+/-388.81 | | | 227.06+/-388.81 | |
| | | | 65.4+/-275.63 | | | 56.3+/-242.23 | |
| | | | N1=40, N2=168 | × | × | N1=40, N2=31 | × |
| | | | Fold Change: 2.9 | | | Fold Change: 3.21 | |
| | | | P-value: .00003 | | | P-value: .0001 | |

| Sed ID | Genbank | Normal vs All | Normal vs Malignant | Normal vs Stage I | Normal vs Stage II | Normal vs Stage III |
|--------|----------|-------------------|---------------------|-------------------|--------------------------------------|---------------------|
| 556 | AI655499 | 46.97+/-125.7 | | | 47.75+/-127.24 | |
| | | 313.1+/-881.21 | | | 453.25+/-957.09 | |
| | | N1=40, N2=168 | × | × | N1=40, N2=31 | × |
| | | Fold Change: 2.28 | | | Fold Change: 2.89 P-value: .00225 | |
| 260 | AI656836 | | | | | 496.58+/-144.86 |
| 3 | | | | | | 242.93+/-91.56 |
| | | × | × | × | × | N1=40, N2=10 |
| | | | | | | Fold Change: 2.07 |
| | | | | | | P-value: .00005 |
| 562 | AI658925 | 545.69+/-343.76 | | | 542.56+/-347.67 | 542,56+/-347.67 |
| | | 259.08+/-213.04 | | | 233.63+/-187.46 | 164.54+/-115.15 |
| | | N1=40, N2=168 | × | × | N1=40, N2=31 | N1=40, N2=10 |
| | | Fold Change: 2.3 | | | Fold Change: 2.42 | Fold Change: 3.62 |
| | | P-value: 0 | | | P-value: .00001 | P-value: .00181 |
| 563 | AI658928 | 230.91+/-89.43 | | | 230.91+/-89.43 | 230.91+/-89.43 |
| } | | 89.62+/-64.08 | | | 75.71+/-63.86 | 130,21+/-102.4 |
| | | N1=39, N2=168 | × | × | N1=39, N2=31 | N1=39, N2=6 |
| | | Fold Change: 2,76 | | | Fold Change: 3.17 | Fold Change: 2.16 |
| | | P-vaine: 0 | | | P-value: 0 | P-value: .02073 |
| 565 | AI659418 | | | | | 261.02+/-116.11 |
| | | | | | | 125.48+/-61.12 |
| | | × | × | × | × | N1=40, N2=10 |
| | | | | | | Fold Change: 2.05 |
| | | | | | | P-value: .00057 |
| 366 | AI659533 | 566.04+/-199.44 | | | 563.4+/-201.34 | 563.4+/-201.34 |
| | | 260.59+/-219.32 | | | 289.88+/-264.96 | 161.05+/-65.87 |
| | | N1=40, N2=168 | × | × | N1=40, N2=31 | N1=40, N2=10 |
| | | Fold Change: 2.68 | | | Fold Change: 2.58 | Fold Change: 3.49 |
| | | P-value: 0 | | | P-value: 0 | P-value: 0 |
| 268 | AI659927 | | | | | 427.88+/-182.76 |
| | | | | | | 161.17+/-86.36 |
| | | × | × | × | × | N1=40, N2=10 |
| | | | | | | Fold Change: 2.71 |
| | | | | | | P-value: .00004 |

| 1 | | | | | 1 | | _ | | Г | | | | T | - | | _ | _ | ٦ | | | | | ٦ | | | | | Т | | | |
|---|---------------------|------------------------------------|---------------|-------------------|-----------------|-------------------|--------------|-------------------|-------------|-----------------|--------------|------------------|----------------|-----------------|-----------------|--------------|------------------|--|----------------|-----------------|--------------|-------------------|-----------------|-----------------|----------------|--------|-------------------|----------------|----------------|--------------|-------------------|
| | Normal vs Stage III | 357.85+/-302.39 | N1=39, N2=6 | Fold Change: 2.63 | P-value: .01212 | | × | | 112+/-77.42 | 229.83+/-111.91 | N1=40, N2=10 | Fold Change: 2.3 | r-vame: .00054 | 172.75+/-104.93 | 326.06+/-163.08 | N1-40, N2-10 | Fold Change: 2.1 | P-value: .00523 | | | × | | | | Þ | < | | | | × | |
| | Normal vs Stage II | | × | | | 1443,34+/-1731.78 | N1=40, N2=31 | Fold Change: 2.82 | | | × | | | | | × | | The second secon | 285.7+/-185.25 | 643.27+/-468.95 | N1=40, N2=31 | Fold Change: 2.17 | P-value: .00012 | 281.86+/-139.36 | 128.45+/-98.58 | D-11 C | Fold Change: 2.34 | P-vame: .00001 | 119.84+/-49.46 | N1=40, N2=31 | Fold Change: 2.13 |
| | Normal vs Stage I | | × | | | | × | | | | × | | | | | × | | | | | × | | | | Þ | < | | | | × | |
| | Normal vs Malignant | | × | | | | × | | | | × | | | | | × | | | | | × | | | | Þ | * | | | | × | |
| | Normal vs All | 119.06+/-130.86 277.49+/-246.97 | N1=39, N2=168 | Fold Change: 2.1 | P-value: 0 | | × | | | | × | | | | | × | | | | | × | | | | Þ | < | | | | × | |
| | Genbank | AI660245 | | | | AI668620 | | | AI671836 | | | | | AI671984 | | | | | AI673539 | | | | | AI673735 | | | | 43004600 | A1674603 | | |
| | Sed ID | 269 | | | | 571 | | | 574 | | | | | 575 | | | | | 579 | | | | | 280 | | | | 202 | 283 | | |
| | # | 288 | | | | 289 | | | 290 | | | | | 291 | | | | | 292 | | | | | 293 | | | | | 294 | | |

| 7 | m wo | Conhant | Mosmol ve All | Normal we Malianant | Normal ve Stone I | Mormal ve Stage II | Wormal ve Stage III |
|-----|--------|----------|---------------------------------|------------------------|--------------------|----------------------|---------------------|
| £ | Sed TD | Gendank | POPULAL VS ALI | TOFILIAL VS PRAIRGIANT | I again as Stage I | Itol mar vs Brage At | TOTAL SO DURES AT |
| 295 | 584 | AI675106 | 353.92+/-130.58 | | | | |
| | | | 100.027-1-101.45 | Þ | Þ | Þ | Þ |
| | | | NI=39, INZ-108 | < | < | 4 | * |
| | | | Fold Change: 2.02 P-value: 0 | | | | |
| 296 | 588 | AI680541 | 516.15+/-202.37 | | | 510.08+/-201.29 | 510.08+/-201.29 |
| | | | 138.66+/-107.52 | | | 149.24+/-129.55 | 101.96+/-86.37 |
| | | | N1=40, N2=168 | × | × | N1=40, N2=31 | N1=40, N2=10 |
| | | | Fold Change: 4.19 | | | Fold Change: 3.77 | Fold Change: 5.63 |
| | | | P-value: 0 | | | P-value: 0 | P-value: .00001 |
| 297 | 290 | AI683036 | 106.33+/-110.74 | | | 106.33+/-110.74 | |
| | | | 328.16+/-267.3 | | | 296.92+/-224.37 | |
| | | | N1=39, N2=168 | × | × | N1=39, N2=31 | × |
| | | | Fold Change: 2.81 | | | Fold Change: 2,63 | |
| | | | P-value: 0 | | ! | P-value: 0 | |
| 867 | 591 | AI683911 | 241,46+/-200.89 | | | 241.46+/-200.89 | 241,46+/-200.89 |
| | | | 35.47+/-57.1 | | | 28.41+/-33.49 | 29,45+/-35.37 |
| | | | N1=40, N2=168 | × | × | N1=40, N2=31 | N1=40, N2=10 |
| | | | Fold Change: 4.64 | | | Fold Change: 5.05 | Fold Change: 5.29 |
| | | | P-value: 0 | | | P-value: 0 | P-value: 0 |
| 562 | 592 | AI684457 | 96.99+/-74.31 | | | | |
| | | | 233.36+/-405.3 | | | | |
| | | | N1=40, N2=168 | × | × | × | × |
| | | | Fold Change: 2 | | | | |
| | | | P-value: .00001 | | | | |
| 300 | 593 | AI686114 | 375.54+/-271.13 | | | 374.48+/-274.59 | |
| | | | 158.93+/-158.15 | | | 155.96+/-124.29 | |
| | | | N1=40, N2=168 | × | × | N1=40, N2=31 | × |
| | | | Fold Change: 2.67 | | | Fold Change: 2.4 | |
| | | | P-value: 0 | | | P-value: .00006 | |
| 301 | 594 | AI686316 | 255.25+/-97.58 | | 255.25+/-97.58 | 255.25+/-97.58 | 255.25+/-97.58 |
| | | | 102.19+/-93.67 | | 120.04+/-93.28 | 79.78+/-67.6 | 51.42+/-78.59 |
| | | | N1=39, N2=168 | × | NI=39, N2=10 | N1=39, N2=31 | N1=39, N2=6 |
| | | | Fold Change: 2.55 | | Fold Change: 2.52 | Fold Change: 2.99 | Fold Change: 3.34 |
| | | | P-value: 0 | | F-value: .0397 | r-vaine: 0 | r-value: .00001 |

| # | Seq ID | Genbank | Normal vs All | Normal vs Malignant | Normal vs Stage I | Normal vs Stage II | Normal vs Stage III |
|-----|--------|----------|---------------------------------|---------------------|-------------------|--------------------|---------------------|
| 302 | 595 | AI689747 | 229.57+/-81.34 | | | 229.57+/-81.34 | 229.57+/-81.34 |
| | | | 113 53+/-98 71 | | | 120.42+/-76.86 | 74.4+/-34.38 |
| | | | N1=39. N2=168 | × | × | N1=39, N2=31 | N1=39, N2=6 |
| | | | Fold Change: 2,2 | | | Fold Change: 2.07 | Fold Change: 2.9 |
| | | | P-value: 0 | | | P-value: 0 | P-value: 0 |
| 303 | 597 | AI691077 | 201.94/-110.51 | | | | |
| | | | 84.62+/-106.12 | | | | |
| | | | N1=40, N2=168 | × | × | × | × |
| | | | Fold Change: 2.35 P-value: 0 | | | | |
| 304 | 200 | AI692687 | | | | | 215.65+/-82.22 |
| | | | | | | | 106.33+/-67.45 |
| | | | × | × | × | × | N1=39, N2=6 |
| | | | | | | | Fold Change: 2.27 |
| | | | | | | | P-value: .00376 |
| 305 | 603 | AI693690 | | | | | 187.55+/-153.71 |
| | | | | | | | 419.84+/-166.89 |
| | | | × | × | × | × | N1=40, N2=10 |
| | | | | | | | Fold Change: 2.84 |
| | | | | | | | P-value: .00002 |
| 306 | 604 | AI694059 | 256.63+/-189.71 | | | 256.63+/-189.71 | 256.63+/-189.71 |
| | | | 63.49+/-68.69 | | | 45.9+/-74.24 | 24.83+/-23.88 |
| | | | N1=39, N2=168 | × | × | N1=39, N2=31 | N1=39, N2=6 |
| | | | Fold Change: 3.4 | | | Fold Change: 3.8 | Fold Change: 5.35 |
| | | 1 | P-value: 0 | | | P-value: 0 | P-value: 0 |
| 307 | 909 | AI695684 | | | | | 284.08+/-85.43 |
| | | | | | | | 138.19+/-31.83 |
| | | | × | × | × | × | N1=40, N2=10 |
| | | | | | | | Fold Change: 2.01 |
| | | | | | | | P-value: 0 |
| 308 | 809 | AI698134 | 801.15+/-394.52 | | 803.86+/-399.3 | 803.86+/-399.3 | 803.864/-399.3 |
| | | | 295.3+/-157.04 | | 305.4+/-156.08 | 238.5+/-98.66 | 290.26+/-225.95 |
| | | | N1=40, N2=168 | × | N1=40, N2=6 | N1=40, N2=31 | N1=40, N2=10 |
| | | | Fold Change: 2.66 | | Fold Change: 2.52 | Fold Change: 3.16 | Fold Change: 3.23 |
| | | | P-value: 0 | | P-value: .00483 | P-value: 0 | P-value: .00182 |

| Seg ID Genbank | Genbank | | Normal vs All | Normal vs Malignant | Normal vs Stage I | Normal vs Stage II | Normal vs Stage III |
|------------------------------|---------------|---------------|---------------|---------------------|-------------------|--------------------|---------------------|
| AT700484 A | | 404 49+/-2 | 04.86 | | 404 49+/-304 86 | 404.49+/-304.86 | 404.49+/-304.86 |
| - COLOG-101- | | 149.98+/-9 | 3.82 | | 151.52+/-60.91 | 139.3+/-55.78 | 132,26+/-50.46 |
| N1=40.N2=168 | N1=40, N2 | N1=40, N2= | =168 | × | N1=40, N2=6 | N1=40, N2=31 | N1=40, N2=10 |
| Fold Change: 2.59 | Fold Change | Fold Change | 2.59 | | Fold Change: 2.49 | Fold Change: 2.66 | Fold Change: 2.75 |
| P-value: 0 | P-value: | P-value: | 0 | | P-value: .00796 | P-value: 0 | P-value: 0 |
| 612 AI701034 | AI701034 | | | | | | 215.78+/-96.65 |
| × | × | × | | × | × | × | N1=40, N2=10 |
| | | | | | | | Fold Change: 2.03 |
| | | | | | | | P-value: .00018 |
| 613 AI703441 | AI703441 | | | | | 166.32+/-110.39 | 166,32+/-110,39 |
| | | | | | | 348.27+/-255.22 | 332,43+/-137.78 |
| × | × | × | | × | × | N1=40, N2=31 | N1=40, N2=10 |
| | | | | | | Fold Change: 2 | Fold Change: 2.31 |
| | | | | | | P-value: .00052 | P-value: .00003 |
| 614 AI703451 | AI703451 | | | | | 58.72+/-154.11 | |
| | | | | | | 479.31+/-977.92 | |
| × | × | × | | × | × | N1=40, N2=31 | × |
| | | | | | | Fold Change: 3.05 | |
| | | | | | | P-value: .00313 | |
| 615 ALT07589 446.78+/-668.95 | | 446.78+/-668. | 25 | | | 414.9+/-646.18 | 414.9+/-646.18 |
| 359.77+/-1435.2 | 359.77+/-143 | 359.77+/-143 | 5.2 | | | 236.79+/-693.96 | -8.82+/-71.63 |
| N1=40, N2=168 | N1=40, N2= | N1=40, N2= | 891 | × | × | N1=40, N2=31 | N1=40, N2=10 |
| Fold Change: 2.73 | Fold Change: | Fold Change: | 2.73 | | | Fold Change: 2.49 | Fold Change: 4.36 |
| P-value: .00099 | P-value: .000 | P-value: .000 | 660 | | | P-value: .01893 | P-value: .00006 |
| 617 AI720763 | AI720763 | | | | | | 216,2+/-123.53 |
| | | | | | | | 94.49+/-39.1 |
| × | × | × | | × | × | × | N1=40, N2=10 |
| | | | | | | | Fold Change: 2.05 |
| | | | | | | | P-value: .00026 |
| 618 AI732274 926.55+/-985.51 | | 926.55+/-985 | 51 | | 947.08+/-989.69 | 947.08+/-989.69 | 947.08+/-989.69 |
| 133.4+/-273.85 | 133.4+/-273 | 133.4+/-273 | .85 | | 92.83+/-139.88 | 17.45+/-85.58 | 25.59+/-138.38 |
| N1=40, N2=168 | N1=40, N2= | N1=40, N2= | 168 | × | N1=40, N2=6 | NI=40, N2=31 | N1=40, N2=10 |
| Fold Change: 7.17 | Fold Change: | Fold Change: | 7.17 | | Fold Change: 8.37 | Fold Change: 13.2 | Fold Change: 14.18 |
| P-vaine: 0 | P-vaine: | P-vaine: | 0 | | F-value: .00451 | r-value: 0 | F-value: 0 |

| # | Seq ID | Genbank | Normal vs All | Normal vs Malignant | Normal vs Stage I | Normal vs Stage II | Normal vs Stage III |
|-----|--------|----------|-------------------|---------------------|-------------------|--------------------|---------------------|
| 316 | 619 | AI733679 | 319.55+/-589.9 | | 325.9+/-596.22 | 325.9+/-596.22 | 325.9+/-596.22 |
| | | | 41 3+/-61 11 | | 26.3+/-20.79 | 23.3+/-13,87 | 22.92+/-14.21 |
| | | | N1=40. N2=168 | × | N1=40, N2=6 | N1=40, N2=31 | N1=40, N2=10 |
| | | | Fold Change: 3.96 | | Fold Change: 4.56 | Fold Change: 5.05 | Fold Change: 5.39 |
| | | | P-value: 0 | | P-value: .00001 | P-value: 0 | P-value: 0 |
| 317 | 621 | AI740483 | | | | | 519.7+/-165.79 |
| | | | | | | | 245.74+/-94.64 |
| | | | × | × | × | × | N1=39, N2=6 |
| | | | | | | | Fold Change: 2.11 |
| | | | | | | | P-value: .00011 |
| 318 | 622 | AI740516 | 23.68+/-49.95 | | | 24.14+/-50.52 | |
| | | | 230.26+/-254.07 | | | 211.05+/-266.38 | |
| | | | N1=40, N2=168 | × | × | N1=40, N2=31 | × |
| | | | Fold Change: 5.25 | | | Fold Change: 4.63 | |
| | | | P-value: 0 | | | P-value: 0 | |
| 310 | 623 | AT740621 | 231.84+/-247.13 | | | 231.84+/-247.13 | · 231.84+/-247.13 |
| | ļ | | 51.4+/-63.67 | | | 55.89+/-78.15 | 58.59+/-71.76 |
| | | | N1=40, N2=168 | × | × | N1=40, N2=31 | N1=40, N2=10 |
| | | | Fold Change: 2.99 | | | Fold Change: 2.92 | Fold Change: 2.89 |
| | | | P-value: 0 | | | P-value: .00006 | P-value: .00645 |
| 320 | 624 | AI741026 | 324.97+/-140.14 | | | 321.67+/-140.4 | |
| | | | 152.41+/-75.46 | | | 156.42+/-93.26 | |
| | | | N1=40, N2=168 | × | × | N1=40, N2=31 | × |
| | | | Fold Change: 2.1 | | | Fold Change: 2.06 | |
| 1 | | | P-vame: 0 | | | r-value; o | 44 400 1 100 110 |
| 371 | 627 | AI/42002 | 109.124/-131.8 | | 111./8+/-152.43 | 111./8+/-132.43 | 111./04/-152.45 |
| | | | 356,63+/-240.05 | | 392.25+/-219.22 | 395.11+/-278.86 | 430.05+/-236.92 |
| | | | N1=40, N2=168 | × | N1=40, N2=6 | N1=40, N2=31 | N1=40, N2=10 |
| | | | Fold Change: 3.82 | | Fold Change: 4.18 | Fold Change: 3.97 | Fold Change: 4.87 |
| | | | P-value: 0 | | P-value: .00524 | P-value: 0 | P-value: 0 |
| 322 | 628 | AT742057 | 200.43+/-229.58 | | | 203.47+/-231.77 | 203.47+/-231.77 |
| | | | 445.89+/-295.68 | | | 460.52+/-275.2 | 448.38+/-232.46 |
| | | | NI=40, N2=168 | × | × | N1=40, N2=31 | N1=40, N2=10 |
| | | | Fold Change: 2.37 | | | Fold Change: 2.6 | Fold Change: 2.46 |
| | | | P-value: 0 | | | P-value: 0 | P-value: .00303 |

| 1 | 9 | Conhank | Mormal vie All | Normal vs Malionant | Normal vs Stage I | Normal vs Stage II | Normal vs Stage III |
|-----|--------|------------|-------------------|---------------------|-------------------|--------------------|---------------------|
| | oed an | CCIIDAIIA | TOTAL STATE | 0 | | 607 86+/-254 44 | 697 86+/-254 44 |
| | 9 | AI750575 | /02:71+7-253:03 | | | 30 000) : 00 200 | 275 214/ 286 22 |
| | | | 412.71+/-328.09 | | | 511.65-1-296.43 | 77.007-117-017 |
| | | | N1=40. N2=168 | × | × | N1=40, N2=31 | N1=40, N2=10 |
| | | | Fold Change: 2 14 | | | Fold Change: 2.36 | Fold Change: 3.89 |
| | | | D-value. 0 | | | P-value: .00005 | P-value: .00699 |
| - 1 | 544 | AT721.420 | 141 95+/-187 15 | | | 144.67+/-188.73 | 144.67+/-188.73 |
| | 140 | 0C#TC/TW | 659 00-1-724 54 | | | 466,36+/-471.12 | 306.05+/-201.86 |
| | | | PC-121-17-00-17-0 | Þ | × | N1=40, N2=31 | N1=40, N2=10 |
| | | | NI=40, NZ=106 | < | • | Fold Change: 3.51 | Fold Change: 2.79 |
| | | | Ford Change: 4.71 | | | P-value: 0 | P-value: .00492 |
| | 45 | VOJUZITA V | 275 26±/, 190 87 | | | 369.82+/-189.17 | 369.82+/-189.17 |
| 337 | 740 | ABUAC/14 | 160 204/ 155 84 | | | 148.94+/-124.57 | 76.67+/-82.11 |
| | | | 891=CN 0V-IN | * | × | N1=40, N2=31 | N1=40, N2=10 |
| | | | D-14 Chance 2 01 | • | | Fold Change; 2.85 | Fold Change: 5.81 |
| | | | Fold Cuange: 5.01 | | | P-value: .00001 | P-value: .00014 |
| | - 1 | A THEODY | 026 NTLL 657 44 | | 833.52+/-665.83 | 833.52+/-665.83 | 833.52+/-665.83 |
| 555 | 643 | A1/304/3 | 152 01-7 456 44 | | 215.9+/-314.77 | 164.76+/-581.67 | 52.12+/-64.89 |
| | | | N1=40 N2=168 | × | N1=40, N2=6 | N1=40, N2=31 | N1=40, N2=10 |
| | | | Fold Change: 9 32 | | Fold Change: 6.2 | Fold Change: 10.99 | Fold Change: 14.24 |
| | | | D-refue. 0 | | P-value: .02859 | P-value: 0 | P-value: 0 |
| | - 1 | 17750400 | O Commany | | | | 383.82+/-126.82 |
| 33 | 4 | AL/58408 | | | | | 188,25+/-213,28 |
| | | | Þ | > | × | × | N1=40, N2=10 |
| | | | < | : | | | Fold Change: 3.58 |
| | | | | | | | P-value: .01039 |
| | 242 | A1760310 | 208 60+/-65 34 | | 208.69+/-65.34 | 208.69+/-65.34 | 208.69+/-65.34 |
| 333 | | WT/00312 | 105 70+/-40 89 | | 102.62+/-62.32 | 102.48+/-38.82 | 104.41+/-52.84 |
| | | | M1=20 M2=168 | * | N1=39, N2=10 | N1=39, N2=31 | N1=39, N2=6 |
| | | | Told Change: 7 | 1 | Fold Change: 2.02 | Fold Change: 2.07 | Fold Change: 2.13 |
| | | | Daralne. | | P-value: .01579 | P-value: 0 | P-value: .00136 |
| 356 | 949 | AT760370 | | | | 691.21+/-512.28 | 691.21+/-512.28 |
| | | 01000170 | | | | 259.02+/-226.94 | 142.47+/-133.11 |
| | | | × | × | × | N1=40, N2=31 | N1=40, N2=10 |
| | | | : | | | Fold Change: 2.3 | Fold Change: 4.03 |
| | | | | | | P-value: .00272 | P-value: .00082 |

| # | Sea ID | Genbank | Normal vs All | Normal vs Malignant | Normal vs Stage I | Normal vs Stage II | Normal vs Stage III |
|-----|--------|----------|-------------------|---------------------|-------------------|---------------------------------|---------------------|
| 337 | 648 | AI760589 | | | | 81.29+/-140.48 | |
| | | | × | × | × | N1=40, N2=31 | × |
| | | | | | | Fold Change: 3.26 P-value: 0 | |
| 338 | 649 | AI761241 | 891.41+/-331.82 | | | 883.3+/-332.12 | 883,3+/-332,12 |
| | | | 417.19+/-273.72 | | | 352.83+/-245.83 | 238,36+/-101.2 |
| | | | N1=40, N2=168 | × | × | N1=40, N2=31 | N1=40, N2=10 |
| | | | Fold Change: 2.32 | | | Fold Change: 2.7 | Fold Change: 3.83 |
| | | | P-value: 0 | | | P-value: 0 | P-value: .00005 |
| 339 | 650 | AI761274 | 346.01+/-181.77 | | | 342.36+/-182.65 | 342,36+/-182.65 |
| | | | 114.5+/-91.9 | | | 121.75+/-92.05 | 77.48+/-76.11 |
| | | | N1=40, N2=168 | × | × | N1=40, N2=31 | N1=40, N2=10 |
| | | | Fold Change: 3.44 | | | Fold Change: 3.23 | Fold Change: 4.73 |
| | | | P-value: 0 | | | P-value: 0 | P-value: .00028 |
| 940 | 651 | AI761782 | | | | | 41.02+/-103.08 |
| | | | | | | | 282,44+/-98.22 |
| | | | × | × | × | × | N1=39, N2=6 |
| | | | | | | | Fold Change: 6.26 |
| | | | | | | | P-value: 0 |
| 341 | 652 | AI761844 | 284.64+/-141.48 | | | 278.83+/-138.41 | 278.83+/-138.41 |
| | | | 117.61+/-150.43 | | | 132.4+/-302.8 | 94.71+/-60.55 |
| | | | N1=40, N2=168 | × | × | N1=40, N2=31 | N1=40, N2=10 |
| | | | Fold Change: 2.8 | | | Fold Change: 3,35 | Fold Change: 3.05 |
| | | | P-value: 0 | | | P-value: 0 | P-value: .0007 |
| 342 | 654 | AI763298 | 247.25+/-99.93 | | | 242.06+/-95.62 | 242.06+/-95.62 |
| | | | 113.86+/-100.53 | | | 148.77+/-173.93 | 84.64+/-67.22 |
| | | | N1=40, N2=168 | × | × | N1=40, N2=31 | N1=40, N2=10 |
| | | | Fold Change: 2.6 | | | Fold Change: 2.29 | Fold Change: 3.32 |
| | | | P-value: 0 | | | P-value: .00002 | P-value: .00045 |
| 343 | 655 | AI766029 | 265.74+/-522.75 | | 271.74+/-528.19 | 271.74+/-528.19 | 271.74+/-528.19 |
| | | | . 15.92+/-35.83 | | 11.19+/-13.38 | 4.73+/-8.61 | 8.83+/-25.01 |
| | | | N1=40, N2=168 | × | N1=40, N2=6 | N1=40, N2=31 | N1=40, N2=10 |
| | | | Fold Change: 3.94 | | Fold Change: 4.33 | Fold Change: 4.64 | Fold Change: 4.09 |
| | | | P-value: 0 | | P-value: 0 | P-value: 0 | P-value: 0 |

| Seq 1D Genbunk Normal vs All Normal vs Milgrant Normal vs Shage II Normal vs Shage II | 1 | | | | | | | |
|--|-----|--------|----------|-------------------|-------------------------------------|-------------------|--|---|
| 4 650 AT7685777 X | × | Seq 1D | Genbank | Normal vs All | Normal va Mali | | | |
| Mathematical Characteristics | 344 | | AI768777 | | THOU THESE AS A PARTICULAR PROPERTY | Normal vs Stage I | Normal vs Stage II | Normal vs Stage III |
| 661 AT709559 X | | | | × | × | × | × | 263.99+/-143.63 686.1+/-326.01 NI=40, N2=10 |
| March Marc | 345 | i | AI769559 | | | | | P-value: .00011 |
| 662 A170060 140.024-101.02 140.024-101.02 Fold Changer 2.79 X 120.24-101.02 Fold Changer 2.14 X 120.24-101.02 Fold Changer 2.14 X 120.24-101.02 Fold Changer 2.14 X X 120.24-101.02 Fold Changer 2.14 X X X X X X X X X | | - 1 | | × | × | × | 213.16+/-92.6 101.34+/-76.51 N1=39, N2=31 Fold Change: 2.13 | × |
| Mi = 40, N2 = 168, Page 2, 19 | 9 | | AI770080 | 450.33+/-335.5 | | | P-value: 0 | |
| No. | | | | 166.16+/-190.29 | | | 453.96+/-339.09 | 453.96+/-339.09 |
| Fold Changes 2.79 | | | | N1=40, N2=168 | Þ | 3 | 129.2+/-107.13 | 94.56+/-93.08 |
| 663 AI788490 TG-Lit-yag-171 663 AI788490 TG-Lit-yag-171 1513.284-246.71 665 AI792405 X X X X 2020.04-224.90 667 AI792405 X X X X 2020.04-224.90 667 AI792817 112.384+443.52 669 AI792817 112.384+443.52 Fold Changer 2.10 Fold Changer 2. | | | | Fold Change: 2.79 | < | × | N1=40, N2=31 | N1=40, N2=10 |
| | 1 | | | P-value: 0 | | | Fold Change: 3.2 | Fold Change: 4.55 |
| 1513-841-5107-1 | | 963 | AI783490 | 767.15+/-947.17 | | | P-value: 0 | P-value: .00046 |
| Name | | | | 1513.38+/-2167.1 | | | | |
| Fold Changer 2.14 | | | | NI=40, N2=168 | Þ | ; | | |
| 665 AI791632 2 Profiles, 00537 X X 667 AI792403 X X X 667 AI792404 X X X 7 Paraline, 0.23 X X X 869 AI792817 112384-143.27 Y Y 869 AI792817 112384-143.27 Y Y 860 AI792817 Fold Change, 2.27 Fold Change, 2.27 860 AI792817 Fold Change, 2.27 Fold Change, 2.27 860 AI792817 Fold Change, 2.27 Fold Change, 3.97 860 AI792817 Fold Change, 3.97 Fold Change, 3.97 860 A | | | | Fold Change; 2,14 | ¢ | × | × | × |
| 665 APP91632 255.664-11,100.68 X X 667 APP92405 X X X 667 APP92405 X X X F-value: 0. P-value: 0. 99.8294-112.78 302.014-224.89 669 AJP92817 112.384-445.32 X X NII-40, N2-31 669 AJP92817 112.384-445.32 X NII-40, N2-31 Fold Change: 2.02 669 AJP92817 112.384-445.32 X NII-40, N2-31 Fold Change: 2.02 Fold Change: 2.07 NII-40, N2-31 AND-40, N2-61 Fold Change: 2.07 Fold Change: 2.07 Fold Change: 3.07 Fold Change: 3.07 Fold Change: 3.07 Fold Change: 3.07 | 1 | | | P-value: ,00537 | | | | |
| 12.58+L46.55 | 00 | 999 | AI791632 | 255.06+/-150.68 | | | | |
| Min = 0, Nin = 0, N | | | | 122.84+/-86.83 | | | | 255.06+/-150.68 |
| Fold Change; 2.2.2 | | | | N1=40, N2=168 | * | ۵ | | 96.82+/-63.86 |
| 667 AI7924045 P-value: 0 95.829-L12.78 X X 202.014-J234.89 K X 10.014-J234.89 Fold Change: 2.02 Fold Change: 2.02 Fold Change: 3.07 112.384-L443.22 Fold Change: 3.07 112.384-L443.22 Fold Change: 3.04 112.384-L443.22 Fold Change: 3.04 112.384-L443.22 Fold Change: 3.07 Fold Change: 3.07 | | | | Fold Change: 2.22 | 4 | × | × | NI=40, N2=10 |
| 667 AP924045 X X 2028-4-112.78 669 AP92481 112.384-443.22 Fold Changer 2.07 67 AP92481 558.024-6937.45 III.2384-143.52 IIII.2384-143.52 III. | - 1 | | | P-value: 0 | | | | Fold Change: 2.58 |
| X X X 202.014-224.89 S25-7-112.28 Fold Changer 2.02 Fold Changer 2.02 Fold Changer 2.03 Fold | | 299 | AI792405 | | | | | P-value: .00084 |
| X | | | | | | | 93.82+/-112.78 | |
| 669 AI792817 112.384-Li43.52 Pod Change 2.102 NI=40, N2=168 NI=40, N2=168 X NI=40, N2=168 Pod Change 3.33 NI=40, N2=31 Pod Change 3.33 NI=40, N2=31 Pod Change 3.39 Pod Change 3.30 Pod Chang | | | | × | × | × | 202.01+/-224.89 | |
| 669 AI792817 112384-i443.52 P-athic 00448 558.024-9327-45 N1-40, N2-468 X N1-40, N2-51 Fold Change; 3.33 P-athice 0 P-athic 00448 Supplementary N2-46 Fold Change; 3.33 Fold Change; 3.37 Fold C | | | | | | | INI-40, NZ=31 | × |
| 669 AI792817 112384-143.52 E-2-2-2-2-2-2-2-2-2-2-2-2-2-2-2-2-2-2- | - 1 | | | | | | Fold Change: 2.02 | |
| X X 510.794-532 510.794-50 | | 699 | AI792817 | 112.38+/-143.52 | | | F-value: .00448 | |
| X X X X X X X X X X X X X X X X X X X | | | | 558.02+/-937.45 | | | 510 701 (557 67 | 112.38+/-143.52 |
| Fold Change: 3.97 Postfine: 0 | | | | N1=40, N2=168 | × | × | MI-40 M2-21 | 849.79+/-711.76 |
| Payalner 0 | | | | Fold Change: 3.33 | | | Fold Change: 2 07 | NI=40, N2=10 |
| | | | | r-value: 0 | | | P-value: 0 | rota Campe: 7.3 |

| | | | | | and the same of | | |
|-----|--------|----------|-------------------|---------------------|-------------------|--------------------|---------------------|
| # | Seq ID | Genbank | Normal vs All | Normal vs Malignant | Normal vs Stage I | Normal vs Stage II | Normal vs Stage III |
| 32 | 0/9 | AI795953 | 407.3+/-278.56 | | 407.3+/-278.56 | 407.3+/-278.56 | 407.3+/-278.56 |
| | | | 146.44+/-111.32 | | 173.14+/-117,81 | 122,61+/-57.78 | 137.64+/-123.78 |
| | | | N1=39 N2=168 | × | NI=39, NZ=10 | N1=39, N2=31 | N1=39, N2=6 |
| | | | Fold Change: 2.62 | | Fold Change: 2.32 | Fold Change: 2.89 | Fold Change: 3.16 |
| | | | P-value: 0 | | P-value: .03224 | P-value: 0 | P-value: .00069 |
| 352 | 671 | AI796083 | | | 114.7+/-66.6 | | |
| | | | | | 207.1+/-77.72 | | |
| | | | × | × | N1=40, N2=6 | × | × |
| | | | | | Fold Change: 2.06 | | |
| 353 | 672 | AT796210 | 300.25+/-171.19 | | | 300,25+/-171,19 | 300.25+/-171.19 |
| | | | 136.42+/-160.54 | | | 142.27+/-164.88 | 68.23+/-32.91 |
| | | | N1=39, N2=168 | × | × | N1=39, N2=31 | N1=39, N2=6 |
| | | | Fold Change: 2.45 | | | Fold Change: 2.36 | Fold Change: 3.93 |
| | | | P-value: 0 | | | P-value: 0 | P-value: 0 |
| 354 | 673 | AI797063 | 217.39+/-201.68 | | | 220.19+/-203.53 | 220.19+/-203.53 |
| | | | 181.18+/-462.43 | | | 161.62+/-454.65 | 58.77+/-92.21 |
| | | | N1=40, N2=168 | × | × | N1=40, N2=31 | N1=40, N2=10 |
| | | | Fold Change: 2.2 | | | Fold Change: 3.09 | Fold Change: 3.3 |
| | | | P-value: .00014 | | | P-value: .00028 | P-value: .00535 |
| 355 | 674 | AIT97276 | 270.66+/-135.07 | | 271.48+/-136.73 | 271.48+/-136.73 | 271.48+/-136.73 |
| | | | 91.42+/-65.13 | | 110.91+/-50.09 | 84.22+/-53.14 | 51.53+/-37.65 |
| | | | N1=40, N2=168 | × | N1=40, N2=6 | N1=40, N2=31 | N1=40, N2=10 |
| | | | Fold Change: 3.1 | | Fold Change: 2.28 | Fold Change: 3.2 | Fold Change: 5.08 |
| | | | P-value: 0 | | P-value: .00368 | P-value: 0 | P-value: 0 |
| 356 | 675 | AI797788 | | | | | 268.73+/-161.42 |
| | | | | | | | 121.99+/-102 |
| | | | × | × | × | × | N1=40, N2=10 |
| | | | | | | | Fold Change: 2.27 |
| 1 | | | | | | - | P-vame: .0082/ |
| 327 | 929 | AI798144 | 74.39+/-112.61 | | | | |
| | | | N1=39, N2=168 | × | × | × | × |
| | | | Fold Change: 2.13 | | | | |
| | | | P-value: 0 | | | | |

| | Seq ID | Genbank | Normal vs All | Normal vs Malignant | Normal vs Stage I | Normal vs Stage II | Normal vs Stage III |
|-------|--------|--------------|--------------------|---------------------|---|--------------------|----------------------|
| 358 (| 8/9 | AI799784 | 599.82+/-379.39 | | 603.99+/-383.42 | 603.99+/-383.42 | 603,99+/-383,42 |
| | | | 67.15+/-81.53 | | 148.43+/-186.96 | 63.08+/-91.29 | 38 2+/-47 49 |
| | | | NI=40 N2=168 | * | N1=40 N2=6 | N=40 N2=31 | N1=40 N2=10 |
| | | | Hold Change: 0.19 | | Bold Changes 5 77 | Pold Change: 10.41 | D-14 Change: 14 10 |
| | | | P-value: 0 | | P-value: 01750 | P-value: 0.+1 | Fold Custings: 14.19 |
| 350 | 189 | ATR01545 | | | (C) | 2 | 107 724/ 63 11 |
| | 1 | CHOOLEGE CO. | | | | | 10///21/-03:11 |
| | | | | | | | 240.16+/-135.66 |
| | | | × | × | × | × | N1=40, N2=10 |
| | | | | | | | Fold Change: 2.33 |
| | | | | | | | P-value: .00036 |
| 360 (| 289 | AI803208 | | | | | 358.64+/-152.22 |
| | | | | | | | 196.65+/-140.61 |
| | | | × | × | × | × | N1=39, N2=6 |
| | | | | 1 | 1 | | Fold Change: 2.05 |
| | | | | | | | P-value: .01446 |
| 361 | 683 | AI803648 | 107.03+/-109.9 | | | | |
| | | | 295.41+/-304.62 | | | | |
| | | | N1=39 N2=168 | × | > | > | > |
| | | | Fold Change: 2.21 | 4 | 4 | < | < |
| | | | P-value: 0 | | | | |
| 269 | 109 | ATOMAGA | 207 221 222 04 | | | 17 700 7170 000 | 10000110000 |
| | 5 | A1004034 | 30 / 254/ - 252.94 | | | 302.974-234.41 | 302.97+7-234.41 |
| | | | 24.74-103.09 | | | /5.144/-30.3/ | 45.174/-28.13 |
| | | | N1=40, N2=168 | × | × | N1=40, N2=31 | N1=40, N2=10 |
| | | | Fold Change: 3.71 | | | Fold Change: 3.81 | Fold Change: 5.95 |
| | | | P-value: 0 | | | P-value: 0 | P-value: 0 |
| 363 | 989 | AI806221 | 206.98+/-125.27 | | 206.98+/-125.27 | 206.98+/-125.27 | 206.98+/-125.27 |
| | | | 57.8+/-39.92 | | 64.87+/-57.18 | 49.38+/-39.25 | 31.17+/-27.12 |
| | | | N1=39, N2=168 | × | N1=39, N2=10 | N1=39, N2=31 | N1=39, N2=6 |
| | | | Fold Change: 3.4 | | Fold Change: 2.98 | Fold Change: 3.82 | Fold Change: 5.54 |
| | | | P-value: 0 | | P-value: .00675 | P-value: 0 | P-value: 0 |
| 364 (| 289 | AI806324 | 214.04+/-130.5 | | | 211.46+/-131.17 | 211.46+/-131.17 |
| | | | 64.16+/-76.08 | | | 48.58+/-64.25 | 21.48+/-26.9 |
| | | | N1=40, N2=168 | × | × | NI=40, N2=31 | N1=40, N2=10 |
| | | | Fold Change: 3.61 | | | Fold Change: 4.36 | Fold Change: 6.44 |
| | | | P-value: 0 | | | P-value-0 | D stoles O |

| * | Seq ID | Genbank | Normal vs All | Normal vs Malignant | Normal vs Stage I | Normal vs Stage II | Normal vs Stage III |
|-----|--------|----------|-------------------|---------------------|--------------------|--------------------|---------------------|
| 365 | 069 | AI809925 | | | | 283.79+/-467.54 | 283.79+/-467.54 |
| | | | | | | 92:39+/-60:68 | 72.84+/-38.61 |
| | | | × | × | × | N1=40, N2=31 | N1=40, N2=10 |
| | | | | | | Fold Change: 2.37 | Fold Change: 3.03 |
| | | | | | | P-value: .00001 | P-value: .00019 |
| 366 | 169 | AI809953 | 383.3+/-186.88 | | | 383.43+/-189.32 | 383.43+/-189.32 |
| | | | 78.06+/-108.22 | | | 59.3+/-98.99 | 19.05+/-40.62 |
| | | | N1=40, N2=168 | × | × | N1=40, N2=31 | N1=40, N2=10 |
| | | | Fold Change: 5.91 | | | Fold Change: 7.01 | Fold Change: 10.5 |
| | | | P-value: 0 | | | P-value: 0 | P-value: 0 |
| 367 | 692 | AI810042 | | | | | 494,45+/-173.8 |
| | | | | | | | 211.09+/-46.06 |
| | | | × | × | × | × | N1=40, N2=10 |
| | | | | | | | Fold Change: 2.24 |
| | | | | | | | P-value: 0 |
| 388 | 693 | AI810266 | 68.69+/-105.28 | | 68.88+/-106.64 | 68.88+/-106.64 | 68.88+/-106.64 |
| | | | 955.73+/-1984.55 | | 1044.03+/-865.49 | 1634.45+/-2196.2 | 521.04+/-362.47 |
| | | | N1=40, N2=168 | × | N1=40, N2=6 | N1=40, N2=31 | N1=40, N2=10 |
| | | | Fold Change: 6.41 | | Fold Change: 16.18 | Fold Change: 13.8 | Fold Change: 9.22 |
| | | | P-value: 0 | | P-value: .00033 | P-value: 0 | P-value: 0 |
| 369 | 694 | AI810764 | | | | 202.16+/-159.83 | |
| | | | × | * | × | N1=40 N2=31 | × |
| | | | : | ; | ; | Fold Change: 2.01 | : |
| | | | | | | P-value: .00293 | |
| 370 | 700 | AI816806 | 551.09+/-313.71 | | | 560.99+/-311.41 | 560.99+/-311.41 |
| | | | 267.19+/-176.02 | | | 242,91+/-135.88 | 221.89+/-110.66 |
| | | | N1=40, N2=168 | × | × | N1=40, N2=31 | N1=40, N2=10 |
| | | | Fold Change: 2.1 | | | Fold Change: 2.3 | Fold Change: 2.4 |
| | | | P-value: 0 | | | P-value: 0 | P-value: .00018 |
| 371 | 701 | AI816835 | 360.85+/-289.77 | | | 360.85+/-289.77 | 360.85+/-289.77 |
| | | | 146.43+/-155.9 | | | 158.38+/-181.37 | 80.71+/-87.22 |
| | | | N1=40, N2=168 | × | × | N1=40, N2=31 | N1=40, N2=10 |
| | | | Fold Change: 2.82 | | | Fold Change: 2.55 | Fold Change: 4.99 |
| | | | P-value: 0 | | | P-value: .00002 | P-value: .00046 |

| | | 241.5+/-87.57 116.59+/-51.83 | N1=40, N2=10 | Fold Change: 2.2 P-value: .003 | 9 | × | 38 | 00 5 | 27 | _ | . × | .39 | | | 1 × | .75 | | | • | 11 N1=40, N2=10 | .82 Fold Change: 4.85 | | | | × | | | 6 | | | 66 Fold Change: 5.96 |
|-----|-------------------------|---------------------------------|--------------|-----------------------------------|----------------|--------------|-------------------|---|-----------------|-----------------|---------------|-------------------|---------------|--------------|--------------|-------------------|-----------------|----------------|-----------------|-----------------|-----------------------|-----------------|-----------------|-----------------|---------------|-------------------|------------|-----------------|----------------|--------------|----------------------|
| N | NOTHIAL VS STARE II | | × | | 879.11+/-582.6 | N1=40, N2=31 | Fold Change: 2.38 | P-value: .0048 | 394.08+/-228.07 | 173.3+/-120.25 | N1=40, N2=31 | Fold Change: 2.39 | 24.28+/-46.39 | 341.95+/-719 | N1=40, N2=31 | Fold Change: 2.75 | P-vaine: .00243 | 63.25+/-126.56 | 468.97+/-447.94 | N1=40, N2=31 | Fold Change: 5.82 | P-value: 0 | - | | × | | | 349.15+/-245.88 | 114.84+/-86.63 | N1=40, N2=31 | Fold Change: 2.66 |
| , | Normal vs Stage I | | × | | | × | | | | | × | | | | × | | | 63.25+/-126.56 | 247.5+/-201.27 | N1=40, N2=6 | Fold Change: 3.53 | P-value: .04505 | | | × | | | | | × | |
| N | INDITION VS INTRIBUTANT | | × | | | × | | | | | × | | | | × | | | | | × | | | | | × | | | | | × | |
| 100 | Normal vs All | | × | | | × | | ,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,, | 396.29+/-225.56 | 181.53+/-192.42 | N1=40, N2=168 | Fold Change: 2.49 | 7 - Ammo- 7 | | × | | | 60.25+/-126.35 | 301.57+/-342.95 | N1=40, N2=168 | Fold Change: 3.51 | P-value: 0 | -117.72+/-96.27 | 255.76+/-788.64 | N1=40, N2=168 | Fold Change: 2.95 | P-value: 0 | | | × | |
| | Сепрапк | AI817448 | | | AI817698 | | | | AI818579 | | | | AT819198 | | | | | AI819340 | | | | | AI820661 | | | | | AI821432 | | | |
| 8 | Sed In | 702 | | | 703 | | | | 90/ | | | | 707 | • | | | | 208 | | | | | 402 | | | | | 711 | | | |
| | # | 372 | | | 373 | | | | 374 | | | | 7,7 | : | _ | | _ | 376 | | _ | _ | | 377 | _ | | | | 378 | _ | | |

| 200 | Genbank | Normal vs All | Normal vs Malignant | Normal vs Stage I | Normal vs Stage II | Normal ve Stene III |
|-----|----------|---|---------------------|------------------------|--------------------|--|
| 712 | AI821472 | 524.32+/-685.96 | | 510 11+/-694 13 | 510 11+/-60/ 13 | The Same of the sa |
| | | 2000 1 1000 | | CI-LOCATION CONTRACTOR | 013-11-1-034-13 | |
| | | 77.700-1-1932.11 | | 10.64+/-179.34 | 83.31+/-629.25 | |
| | | N1=40, N2=168 | × | N1=40, N2=6 | N1=40, N2=31 | × |
| | | Fold Change: 5.78 | | Fold Change: 4.41 | Fold Change: 5 76 | 1 |
| | | P-value: 0 | | P-value: .02894 | P-value: 0 | |
| 713 | AI823572 | 231.34+/-193.19 | | | 232.21+/-195.63 | |
| | | 124.55+/-179.13 | | | 80 48+/-58 66 | |
| | | N1=40, N2=168 | × | × | N1=40 N2=31 | > |
| | | Fold Change: 2.37 | | | Fold Change: 2 82 | 4 |
| | | P-value: 0 | | | P-value: 0 | |
| 714 | AI823649 | 88.12+/-78.99 | | | 88.12+/-78.99 | 88 17+/-78 00 |
| | | 225,47+/-213.09 | | | 223.88+/-174.27 | 239 23+/-237 02 |
| | | N1=39, N2=168 | × | × | NI=39. N2=31 | N=39 N2=6 |
| | | Fold Change: 2.16 | | | Fold Change: 231 | Fold Change: 22 |
| | | P-value: 0 | | | P-value: 0 | Pavalne: 01300 |
| 720 | AI825877 | | | | 418 53+1/162 52 | A18 52±/ 169 59 |
| | | | | | 174 35+/-58 42 | 183 594/-84 31 |
| | | × | × | × | N1=40 N2=31 | NI-do NO-10 |
| | | | | | Fold Changes 2.20 | T-14 CH 142 |
| | | | | | Ford Change: 2.25 | Fold Change: 2.23 |
| 200 | TOTOLY | 40 000000000000000000000000000000000000 | | | r-value: 0 | r-value: .00011 |
| 77 | A102043/ | 241 17+1.52 | | | 45.86+/-118.99 | |
| | | NI-40 NO-120 | Þ | ļ | 67.100-1-10 | |
| | | T-11 CT 0 0 00 | < | * | N1=40, N2=31 | × |
| | | Fold Change: 2.29 | | | Fold Change: 3.07 | |
| 723 | AI827230 | 711.88+/-268 | | | F-value: .00134 | E |
| | | 275 604/ 167 25 | | | 100.5041-2/0.5/ | /08.38+/-2/0.5/ |
| | | 22.02-1-101.53 | ; | | 266.08+/-114.65 | 273.56+/-97.64 |
| | | NI=40, NZ=168 | × | × | N1=40, N2=31 | NI=40, NZ=10 |
| | | Fold Change: 2.22 | | | Fold Change; 2,67 | Fold Change: 2.5 |
| | | P-value: 0 | | | P-value: 0 | P-value: 0 |
| 724 | AI827248 | | | | | 1009.54+/-892.35 |
| | | ; | i | | | 424.88+/-561.45 |
| | | × | × | × | × | N1=40, N2=10 |
| | | | | | | Fold Change: 3.8 |
| | | | | | | P-value: 0309 |

| | 1 | Seq ID | Genbank | Normal vs All | Normal vs Malignant | Normal vs Stage I | Normal vs Stage II | Normal vs Stage III |
|--|---|--------|----------|--|---------------------|-------------------|--|---|
| AIRSP\$20 72.574-66.55 AIRS\$3102 | | 725 | AJ828075 | × | x | × | × | 55.74+/-25.88 208.73+/-146.52 N1=40, N2=10 Fold Change: 3.44 P-value: .0001 |
| A1857102 X X X A1857788 X X X A185786 X X X A1859620 47774:100.9 A1859620 135.294;308.39 N1959, N2=168 Pold Change: 4.1 Pold Change: 4.1 A1860012 X X | | 730 | AI829520 | 78.954/-63.55 221.754/-308.11 N1=39, N2=168 Fold Change: 2.05 P-value: 0 | × | × | × | × |
| A1857788 X X X A1857856 X X X A1859620 4:774+100.9 A1859620 4:777+100.9 A1859620 15:70-108.39 A17-70-108.39 A17-70 | • | 733 | AI833102 | × | × | × | × | 152.23+/-73.43 301.22+/-134.13 N1=40, N2=10 Fold Change: 2.05 P-value: .00005 |
| A18859620 X X X A18859620 47.74-1.00.9 A1869020 47.74-1.00.9 N18-95-72-166 X Fold Change: 4.1 P-value: 0 X X X | | 734 | A1857788 | X | × | × | × | 201.77+/-88.15 79.71+/-43.65 N1=40, N2=10 Fold Change: 2.61 P-value: ,00082 |
| AIRS9620 4777+100.9 335.294-308.99 N1-93, X2=168 Fold Clunger 4.1 F-value 0 X X X | - | 735 | AI857856 | × | × | × | 157.81+/-89.37 341.48+/-151.28 N1=39, N2=31 Fold Change: 2.22 P-value: 0 | 157.81+/-89.37 329.17+/-128.25 N1=39, N2=6 Fold Change: 2.15 P-value: .00013 |
| AI860012 X X | | 739 | AI859620 | 47.7+/-100.9 335.29+/-308.39 N1=39, N2=168 Fold Change: 4.1 P-value: 0 | × | × | 47.7+/-100.9 292.66+/-218.11 N1=39, N2=31 Fold Change: 4.09 P-value: 0 | 47.74/-100.9 289.43+/-181.71 N1=39, N2=6 Fold Change: 3.84 P-value: .00203 |
| | | 740 | AI860012 | X | × | × | 91.22+/-72.75 226.82+/-105.84 N1=39, N2=31 Fold Change: 2.64 P-value: 0 | 91.22+/-72.75 232.58+/-120.25 N1=39, N2=6 Fold Change: 2.58 P-value: .00185 |

| | | | | | The second second second second | | |
|-----|--------|----------|-------------------|---------------------|------------------------------------|--|---------------------|
| * | Seq ID | Genbank | Normal vs All | Normal vs Malignant | Normal vs Stage I | Normal vs Stage II | Normal vs Stage III |
| 393 | 742 | AI860651 | | | | 183.59+/-123.14 449.58+/-384.81 | |
| | | | × | × | × | N1=40, N2=31 Fold Change: 2.2 P-value: 0 | × |
| 394 | 743 | AI863166 | | | 225.98+/-111.31 | | |
| | | | × | × | N1=39, N2=10 | × | × |
| | | | | | Fold Change: 2.4 P-value: .0473 | | |
| 395 | 747 | AI864898 | 406.1+/-256.59 | | | 401.86+/-258.51 | 401.86+/-258.51 |
| | | | 54.76+/-81.97 | | | 39.21+/-56.87 | 31.77+/-56.32 |
| | | | N1=40, N2=168 | × | × | N1=40, N2=31 | N1=40, N2=10 |
| | | | Fold Change: 8.17 | | | Fold Change: 9.5 | Fold Change: 10.87 |
| | | | P-value: 0 | | | P-value: 0 | P-value: 0 |
| 396 | 748 | AI868289 | | | | 386.08+/-242.58 | 386.08+/-242.58 |
| | | | | | | 184.36+/-111.4 | 156.1+/-103.22 |
| | | | × | × | × | N1~40, N2~31 | N1-40, N2-10 |
| | | | | | | Fold Change: 2.22 | Fold Change: 2.54 |
| | | | | | | P-value: .00003 | P-value: .00207 |
| 397 | 750 | AI871044 | 777.08+/-499.12 | | | 766.39+/-500.99 | 766.39+/-500.99 |
| | | | 185.08+/-242.68 | | | 180.1+/-276.48 | 112.03+/-106.51 |
| | | | N1=40, N2=168 | × | × | N1=40, N2=31 | N1=40, N2=10 |
| | | | Fold Change: 4.86 | | | Fold Change: 5.2 | Fold Change: 6.31 |
| | | | P-value: 0 | | | P-value: 0 | P-value: 0 |
| 368 | 751 | AI872267 | 267.23+/-203.1 | | | 267.23+/-203.1 | 267.23+/-203.1 |
| | | | 574.94+/-319.02 | | | 557.03+/-278.09 | 504.87+/-295.04 |
| | | | N1=40, N2=168 | × | × | N1=40, N2=31 | N1=40, N2=10 |
| | | | Fold Change: 2,41 | | | Fold Change: 2.36 | Fold Change: 2.17 |
| | | | P-value: 0 | | | P-value: .00001 | P-value: .00277 |
| 39 | 753 | AI885164 | | | | | 99.32+/-89.64 |
| | | | | | | | 276.5+/-111.42 |
| | | | × | × | × | × | N1=40, N2=10 |
| | | | | | | | Fold Change: 2.94 |
| | | | | | | | P-value: .00002 |

| # | Seg ID | Genbank | Normal vs All | Normal vs Malignant | Normal vs Stage I | Normal vs Stage II | Normal vs Stage III |
|-----|--------|----------|---|---------------------|---|--|--|
| 900 | 754 | AI885498 | 220.94+/-153.32 574.07+/-702.09 N1=40, N2=168 Fold Change: 2.02 P-value: .00005 | × | × | × | × |
| 101 | 755 | AI885781 | × | × | × | 219.35+/-109.49 556.06+/-397.8 N1=40, N2=31 Fold Change: 2.14 P-value: .00015 | 219.35+/-109.49 582.43+/-483.5 N1=40, N2=10 Fold Change: 2.47 P-value: 00109 |
| 402 | 757 | AI887362 | x | × | × | 817.12+/-289.64 355.42+/-140.94 N1=40, N2=31 Fold Change: 2.26 P-value: 0 | 817.12+/289.64 243.89+/-105.11 N1=40, N2=10 Fold Change: 3.29 P-value: 0 |
| 403 | 758 | AI888322 | × | × | × | 319.22+/-320.74 161.88+/-221.65 N1=40, N2=31 Fold Change: 2.73 P-value: .00024 | 319.22+/-320.74 108.01+/-133 N1=40, N2=10 Fold Change: 3.9 P-value: .00657 |
| 404 | 761 | AI889178 | × | × | 372.23+/-146.77 183.45+/-70.01 N1=59, N2=10 Fold Change: 2.03 P-value: .00518 | 372.23+/-146.77 196.21+/-126.54 N1=39, N2=31 Fold Change: 2.04 Parahor 0 | 372.23+/-146.77 194.59+/-97.59 N1=59, N2=6 Fold Change: 2.03 Parabase: 0.0336 |
| 405 | 762 | AI889959 | × | × | × | 140.79+/-151.42 298.84+/-296.55 N1-40, N2=31 Fold Change: 2.11 P-value: .00028 | 140.79+/-151.42 319.25+/-177.44 N1=40, N2=10 Fold Change: 2.62 P-value: .03302 |
| 406 | 763 | AI890418 | 218.25+/-140.58 37.18+/-39.2 N1=39, N2=168 Fold Change: 3.82 P-value: 0 | × | 218.25+/140.58 76.35+/-66.16 N1=39, N2=10 Fold Change: 3.02 P-value: .02078 | 218.25+/-140.58 35.49+/-32.04 N1=39, N2=31 Fold Change: 3.92 P-value: 0 | 218.25+/-140.58 26.25+/-48 N1=39, N2=6 Fold Change: 4.12 P-value: 0 |
| | | | | | | | |

| # | Seq ID | Genbank | Normal vs All | Normal vs Malignant | Normal vs Stage I | Normal vs Stage II | Normal vs Stage III |
|-----|--------|-----------|-------------------|---------------------|-------------------|--------------------|---------------------|
| 404 | 764 | AI890488 | | | | | 498.18+/-173.26 |
| | | | × | × | × | × | N1=40. N2=10 |
| | | | ! | 1 | ! | ! | Fold Change: 2.19 |
| | | | | | | | P-value: .00095 |
| 408 | 169 | AI912772 | | | | | 124.98+/-38.73 |
| | | | | | | | 344.88+/-151.12 |
| | | | × | × | × | × | N1=40, N2=10 |
| | | | | | | | Fold Change: 2.65 |
| | | | | | | | P-value: .00005 |
| 409 | 772 | AI916544 | 150.68+/-161.18 | | | 151.27+/-163.24 | 151.27+/-163.24 |
| | | | 440.12+/-478.52 | | | 548.66+/-436.19 | 636.35+/-560.17 |
| | | | N1=40, N2=168 | × | × | N1=40, N2=31 | N1=40, N2=10 |
| | | | Fold Change: 2.97 | | | Fold Change: 3.81 | Fold Change: 3.69 |
| | | | P-value: 0 | | | P-value: 0 | P-value: .01086 |
| 410 | 775 | AI917901 | 591.38+/-804.54 | | 601.53+/-812.45 | 601.53+/-812.45 | 601.53+/-812.45 |
| _ | | | 76.3+/-209.26 | | 57.95+/-67.95 | 32.64+/-44.74 | 15.05+/-18.69 |
| | | | N1=40, N2=168 | × | N1=40, N2=6 | N1=40, N2=31 | N1=40, N2=10 |
| | | | Fold Change: 5.07 | | Fold Change: 4.77 | Fold Change: 6.97 | Fold Change: 9.62 |
| _ | | | P-value: 0 | | P-value: .00228 | P-value: 0 | P-value: 0 |
| 411 | 276 | AI921685 | 102.15+/-191.42 | | | 104.01+/-193.56 | |
| _ | | | 374.73+/-772,23 | | | 633.55+/-1087.03 | |
| _ | | | N1=40, N2=168 | × | X | N1=40, N2=31 | × |
| | | | Fold Change: 2.03 | | | Fold Change: 3.3 | |
| 5 | 444 | A T077907 | r-vaine: .0025 | | | 202 23±/ 00 22 | 203 227/ 00 22 |
| ! | | - | | | | 481.16+/-300.69 | 408.30+/-402.89 |
| | | | × | × | × | N1=39, N2=31 | N1=39, N2=6 |
| | | | | | | Fold Change: 2.13 | Fold Change: 2.02 |
| | | | | | | P-value: 0 | P-value: .02385 |
| 413 | 778 | AI923108 | | | | 245.08+/-135.84 | |
| | | | | | | 576.66+/-441.15 | |
| | | | × | × | × | N1=40, N2=31 | × |
| | | | | | | Fold Change: 2.16 | |
| | | | | | | P-value: 0 | |

| # Seq ID Genbank Normal vs Miles | 1 | | | | | | | |
|--|-----|--------|--|-------------------|---------------------|-------------------|--------------------|---------------------|
| 779 AD244028 X | * | Seq ID | Genbank | Normal vs All | Normal vs Malignant | Normal vs Stage I | Normal vs Stage II | Normal vs Stage III |
| National Color National Color | 4 | 419 | AJ924028 | | | | | 426.33+/-126.02 |
| 780 A1924465 448.27#-478.27 448.27#-478.27 781 A1924794 X X X X X X X X X X X X X X X X X X X | | | | Þ | Þ | Þ | : | 190+/-63.01 |
| 780 AJ924465 448.274-478.27 448.274-478.27 781 10.324-433.45 X 10.324-43.45 X 10.324-43.4 | | | | < | < | ٧ | 4 | NI=40, NZ=10 |
| 780 AI924465 448.27+478.27 448.27+478.27 10.234-52.21 10.234-53.45 10.234-52.21 10.234-53.45 10.234-52.21 10.234-53.45 10.234-52.21 10.234-53.45 10.234-53.45 10.234-53.45 10.234-53.45 10.234-53.45 10.234-53.45 10.234-53.15 10.234-53.45 10.234-53.16 10.234-53.45 10.234-53.16 10.234-53.45 10.234-53.16 10.234-53.45 10.234-53.16 10.234-53.16 10.234-53.16 10.234-534-534-534-534-534-534-534-534-534-5 | | | | | | | | Fold Change: 2,25 |
| 12.26+4.1221 11.26+4.1231 11.624+6.38.42 | 14 | 1 | ATOTAKE | 74 974 1±17 944 | | 240 PTL 4TO PT | 240 271 / 470 24 | 140 071 / 470 07 |
| Ni = 40, N | 3 | | COLLANS OF THE PARTY OF THE PAR | 178/1-/- 1780- | | 17.0/1-/17.044 | 17.0/1-/17.0++ | 17.9/4/7/2044 |
| Nu-0, Nu-0 | | | | 123.26+/-122.11 | | 110.32+/-53.45 | 104.34+/-104.06 | 46.51+/-27.28 |
| 781 AD24794 P-value: 0.18 Fold Change: 2.13 782 AD24794 X X X X 784 AD28296 X X X X X 784 AD28296 X X X X X 785 AD28399 302.494-122.77 785 AD28399 302.494-412.77 787 AD34364 215.594-433.77 7887 AD34364 215.594-433.77 7887 AD34364 215.594-433.77 7887 AD34364 215.594-433.77 7887 AD34366 215.594-433.77 7887 AD34366 215.594-433.77 7887 AD34366 215.594-433.77 7887 AD34368 218.594-623.77 7888 AD34368 218.594-623.77 7888 AD34368 218.594-623.77 7889 AD34568 218 | | | | N1=40, N2=168 | × | NI=40, N2=6 | N1=40, N2=31 | N1=40, N2=10 |
| 781 AJ934794 P-value: 0 P-value. 00317 782 AJ93789 X X X X 784 AJ938296 X X X X X 785 AJ92839 302.494-122.77 785 AJ92839 302.494-122.77 787 AJ9436 215.994-7431 | | | | Fold Change: 3.18 | | Fold Change: 2.73 | Fold Change: 3.88 | Fold Change: 6.5 |
| 781 AD24794 X X X X X X 77695 782 AU27695 X X X X X X X 7769 784 AU27899 302.46+1122.77 785 AU27899 302.46+1122.77 786 AU27899 150544-112.77 787 AU27899 150544-112.77 787 AU27899 150544-112.77 788 AU27899 150544-112.77 789 AU27899 | | | | P-value: 0 | | P-value; .00317 | P-value: 0 | P-value: 0 |
| 782 AJ92695 X X X X X X X 784 AJ92695 X X X X X X X X X X X X X X X X X X X | 10 | | AI924794 | | | | 132.27+/-116.84 | 132.27+/-116.84 |
| 782 AJ92695 X X X X X X 7 784 AJ92695 X X X X X X 7 784 AJ926296 X X X X X X X 7 785 AJ926299 300.49+1,122.77 | | | | | | | 324.59+/-170.2 | 321.46+/-186.7 |
| 782 AJ927695 X X X X X X X X X 784 AJ928296 X X X X X X X X X X X X X X X X X X X | | | | × | × | × | N1=39, N2=31 | N1=39, N2=6 |
| 782 AI927695 X X X X X X X 7 784 AI926296 X X X X X X X 7 784 AI926296 X X X X X X X X X 7 785 AI926293 300.49+112.77 7 787 AI934361 21.59+413.27 7 787 AI934361 21.59+414.27 7 787 AI934361 21.59+414.27 7 787 AI94.23.18 78446.2.7 7 41.98+7.26.36 Fold Change: 2.13 Fold Change: 2.18 F | | | | | | | Fold Change: 2.6 | Fold Change: 2.5 |
| 782 AD21695 X X X X X X X 784 AD216296 X X X X X X X X X X X X X X X X X X X | | | | | | | P-value: 0 | P-value: .0029 |
| 784 AD26296 X X X X X X 7 784 AD26296 X X X X X X 7 785 AD26296 X X X X X X X X X X X X X X X X X X X | 11 | 782 | AI927695 | | | | | 624.33+/-219.53 |
| 784 AVD28396 X X X X X X X X X X 784 AVD28396 302.494-122.77 785 AVD28393 302.494-122.77 | | | | | | | | 287.2+/-131.6 |
| 784 AD23296 X X X X X X X X X X X 15.05.25.25.25.25.25.25.25.25.25.25.25.25.25 | | | | × | × | × | × | N1=40, N2=10 |
| 784 AID28296 X X X X X X X 784 AID28299 302.494-122.77 | | | | | | | | Fold Change: 2.22 |
| 784 AD28296 X X X X X X X X 7 1 | | - 1 | | | | | | P-value: .00029 |
| 785 AD54399 302.49+4.122.77 X X X X X 789 AD54399 302.49+4.122.77 | 8 | | AI928296 | | | | | 146.02+/-113.56 |
| 785 AP26893 302.494-122.77 787 AD94364 215.994-787 | | | | | | | | 281.82+/-97.95 |
| 785 AI928393 300.4694-112.77 X | | | | × | × | × | × | N1=40, N2=10 |
| 785 AP28393 302.494-122.77 787 AD94364 215.994-741.37 788 AD943631 7894-741.37 789 AD94363 7894-741.37 789 AD94364 7894-741.37 78 | | | | | | | | Fold Change: 2.05 |
| 785 AP28393 30.049-1/12.77 787 AP34361 21.949-1/2.77 787 AP34361 21.949-1/2.37 787 AP34361 21.949-1/2.37 787 AP34361 X X X 787 AP34361 X X X X X X X X X X X X X X X X X X X | | - 1 | | | | | | P-value: .00016 |
| 166,83+4-119.33 | 6 | | AI928393 | 302.49+/-122.77 | | | 297.26+/-119.77 | 297.26+/-119.77 |
| Ni=d0, N2=168 | | | | 166.83+/-119.33 | | | 142.57+/-77.64 | 130.31+/-96.66 |
| Fold Change: 2/3 F.v.aline: 0 | | | | N1=40, N2=168 | × | × | N1=40, N2=31 | NI=40, N2=10 |
| 787 AU94361 215.994-244.37 220.01+4-245.16 49.54-62.77 47.28+-56.36 NI=d0, N2=16 Fold Camper 3.48 Fold Champer 3.18 P-valine: 0 P-valine: | | | | Fold Change: 2.03 | | | Fold Change: 2.17 | Fold Change: 2.67 |
| 787 AB34361 212-99-4241.37 220-14-243.16 49.3+4-62.77 47.28+-26.36 Ni=40, N2=168 X Ni=40, N2=6 Fold Camper 3.18 Fold Camper 3.18 P-value: 0 P-value: 0.07079 | - 1 | | | P-value: 0 | | | P-value: 0 | P-value: .0034 |
| 4728+/-26,36 X NI-60, N2-6 Fold Changes 318 P-value, :0079 | 20 | 787 | AI934361 | 215.99+/-241.37 | | 220.01+/-243.16 | 220.01+/-243.16 | 220.01+/-243.16 |
| X N1=40, N2=6 Fold Change: 3.18 P-value: .00079 | | | | 49.3+/-62.57 | | 47.28+/-26.36 | 33.19+/-18.11 | 32.04+/-24.58 |
| Fold Change: 3.18 P-value: .00079 | | | | N1=40, N2=168 | × | N1=40, N2=6 | N1=40, N2=31 | N1=40, N2=10 |
| P-value: .00079 | | | | Fold Change: 3.48 | | Fold Change: 3.18 | Fold Change: 4.31 | Fold Change: 4.29 |
| | | | | P-value: 0 | | P-value: .00079 | P-value: 0 | P-value: 0 |

| # | Seq ID | Genbank | Normal vs All | Normal vs Malignant | Normal vs Stage I | Normal vs Stage II | Normal vs Stage III |
|-----|--------|----------|-------------------|---------------------|-------------------|--------------------|---------------------|
| 421 | 788 | AI934407 | | | | | 168.61+/-198.83 |
| | | | ; | \$ | * | \$ | 377.54+/-147.2 |
| | | | 4 | < | 4 | ۷ | NI=40, NZ=10 |
| | | | | | | | P-value: 00001 |
| 422 | 792 | AI935915 | 26.52+/-106.71 | | | 28.01+/-107.68 | 28.01+/-107.68 |
| | | | 267.11+/-334.72 | | | 364.49+/-370.15 | 510.1+/-498.57 |
| | | | N1=40, N2=168 | × | × | N1=40, N2=31 | N1=40, N2=10 |
| | | | Fold Change: 4.41 | | | Fold Change: 6.74 | Fold Change: 8.29 |
| | | | P-value: 0 | | | P-value: 0 | P-value: .00153 |
| 423 | 793 | AI936699 | | | | 769.05+/-392.56 | 769.05+/-392.56 |
| | | | | | | 344.85+/-187.03 | 208,46+/-65,42 |
| | | | × | × | × | N1=40, N2=31 | N1=40, N2=10 |
| | | | | | | Fold Change: 2.04 | Fold Change: 3.07 |
| | | | | | | P-value: .00007 | P-value: 0 |
| 424 | 794 | AI936823 | | | | | 103.06+/-86.01 |
| | | | | | | | 210.69+/-119.04 |
| | | | × | × | × | × | N1=40, N2=10 |
| | | | | | | | Fold Change: 2.33 |
| | | | | | | | P-value: .00059 |
| 425 | 795 | AI937060 | 73.53+/-63.87 | | | 73.53+/-63.87 | 73.53+/-63.87 |
| | | | 221.63+/-200.57 | | | 275.58+/-253.28 | 293.86+/-212.3 |
| | | | N1=39, N2=168 | × | × | N1=39, N2=31 | N1=39, N2=6 |
| | | | Fold Change: 2,37 | | | Fold Change: 2.86 | Fold Change: 3.17 |
| | | | P-value: 0 | | | P-value: 0 | P-value: .00776 |
| 456 | 796 | AI937365 | 458.68+/-248.47 | | 461.09+/-251.24 | 461.09+/-251.24 | |
| | | | 1357.16+/-1303.29 | | 2636.53+/-3163.86 | 1444,29+/-1485.46 | |
| | | | N1=40, N2=168 | × | N1=40, N2=6 | N1=40, N2=31 | × |
| | | | Fold Change: 2.31 | | Fold Change: 3.81 | Fold Change: 2,32 | |
| | | | P-value: 0 | | P-value: .03322 | P-value: .00007 | |
| 427 | 798 | AI939507 | | | | | 67.26+/-38.98 |
| | | | × | × | × | × | N1=40, N2=10 |
| | | | | | | | Fold Change: 2.91 |
| | | | | | | | P-value: .00102 |

| 428 | - | Genbank | Normal vs All | Normal vs Malignant | Normal vs Stage I | Normal ve Stage II | Normal ve Stone III |
|-----|-----|-----------|-------------------|---------------------|-------------------|--------------------|-----------------------|
| | 803 | AI950023 | | | - | Total value of | THOUTING AS OF SELECT |
| | | | | | | 555.397/-291.3 | 535.594/-291.5 |
| | | | ; | i | | 129.73+/-80.19 | 94,45+/-69.77 |
| | | | × | × | × | N1=40, N2=31 | N1-40, N2-10 |
| | | | | | | Fold Change: 2.04 | Fold Change: 2.9 |
| | | | | | | P-value: .00084 | P-value: .00112 |
| 429 | 802 | A1952965 | | | | 161.89+/-108.6 | 161,89+/-108.6 |
| | | | | | | 347 0147 150 00 | 70 000 1 551 |
| | | | > | Þ | Þ | 24.707.710.710 | 401.03+/-202.2/ |
| | | | 4 | < | × | N1=40, N2=31 | N1=40, N2=10 |
| | | | | | | Fold Change: 2.38 | Fold Change: 2.78 |
| П | 200 | | | | | P-value: 0 | P-value: .00455 |
| 5 | 900 | AL953053 | | | | | 96.66+/-61.58 |
| | | | 1 | | | | 403.41+/-323.73 |
| | | | × | × | × | × | N1=40, N2=10 |
| | | | | | | | Fold Change: 3 33 |
| - 1 | | | | | | | P-value: 00265 |
| 431 | 808 | AI954874 | | | 209.46+/-107.86 | | |
| | | | | | 96.19+/-23.42 | | |
| | | | × | × | N1=40, N2=6 | × | × |
| | | | | | Fold Change: 2.01 | | |
| | | | | | P-value: .00017 | | |
| 432 | 810 | AI961206 | | | | 46.06+/-77.63 | 46 06±/_77 63 |
| | | | | | | 204 28 17 400 20 | 00.1/-/100.04 |
| | | | ٠ | Þ | , | 204,287/-102,33 | 225.2/+/-103.4 |
| | | | 4 | < | × | N1=40, N2=31 | N1=40, N2=10 |
| | | | | | | Fold Change: 3.96 | Fold Change: 4.13 |
| 1 | 110 | 4 70 0000 | | | | P-value: 0 | P-value: .00043 |
| ş | /10 | AL9083/9 | 296./1+/-383.1 | | | 295.46+/-388.02 | 295.46+/-388.02 |
| | | | 43.24+/-243.3 | | | 104.96+/-424.91 | -12.25+/-22.65 |
| | | | NI=40, N2=168 | × | × | N1=40, N2=31 | N1=40, N2=10 |
| | | | Fold Change: 4.99 | | | Fold Change: 4.47 | Fold Change: 6.3 |
| - [| | | P-value: 0 | | | P-value: .00001 | P-value: 0 |
| 434 | 818 | AI968904 | 744.48+/-291.11 | | | 738.79+/-292.65 | 2 |
| | | | 370.58+/-143.78 | | | 373.44+/-151.46 | |
| | | | N1=40, N2=168 | × | × | N1=40. N2=31 | > |
| | | | Fold Change: 2 | | | Fold Change: 2.01 | : |
| 1 | | | F-value: 0 | | | P-value: 0 | |

| 1 | | | | | - | | |
|-----|------|----------|-------------------|---------------------|-------------------|----------------------|------------------------|
| # | -1 | Genbank | Normal vs All | Normal vs Malignant | Normal vs Stage I | Normal ve Stone II | Normal ve Otage III |
| 435 | 822 | AI970898 | - 505.69+/-422.11 | | 200 201 / 22 003 | trolling vs Brago II | AND HIST AS STARGE III |
| | | | 140001000 | | 20.9724/-470.92 | 509.53+/-426.92 | 509.53+/-426.92 |
| | | | 142.34+7-84.59 | | 155.46+/-70.61 | 116,99+/-60.55 | 117 04+/-63 48 |
| | | | N1=40, N2=168 | × | N1=40 N2=6 | N1=40 N2=21 | NT-40 NO-10 |
| | | | Fold Change: 3 33 | | D-14 CF | 10-747 OF 147 | NI-40, NZ=10 |
| | | | P-walner 0 | | rotd Change: 2.81 | Fold Change: 3.87 | Fold Change: 3.88 |
| 436 | 823 | AT071441 | 1-value, 0 | | P-value: .00097 | P-value: 0 | P-value: 0 |
| 2 | | 74477 | | | 224.36+/-258.15 | 224.36+/-258.15 | 224.36+/-258.15 |
| | | | | | 25.49+/-91 | 44.4+/-132.79 | -63 46+/-149 66 |
| | | | × | × | N1=40. N2=6 | N1=40 N2=31 | N1-40 N0-10 |
| | | | | | 7-14 0 | 10 711 61 11 | MI-40, INZ-10 |
| | | | | | Fold Change: 5.79 | Fold Change: 2.75 | Fold Change: 4.45 |
| 437 | 7.08 | AT07101A | | | F-value: .01447 | P-value: .0003 | P-value: .0004 |
| • | | | | | | | 121.63+/-64.27 |
| | | | Þ | i | | | 329.2+/-298.85 |
| | | | < | * | × | × | N1=40, N2=10 |
| | | | | | | | Fold Change: 2 35 |
| | - 1 | | | | | | Tota Cuange: 2.23 |
| 438 | 830 | AI972498 | 285 82+/-111 27 | | | | F-vame: .00832 |
| | | | 134 20+7 71 63 | | | 286.51+/-112.64 | 286.51+/-112.64 |
| | | | MI-40 NP-169 | ; | | 124.7+/-61.16 | 109.37+/-47.71 |
| | | | 71. 40, INZ-108 | * | × | N1=40, N2=31 | N1=40, N2=10 |
| | | | roud Change: 2.22 | | | Fold Change: 2,3 | Fold Change: 2.7 |
| 15 | | | P-value: 0 | | | P-value: 0 | P-value: 00057 |
| 5 | 831 | AL972661 | | | | 482 12+/-600 67 | |
| | | | | | | 127 04/ 200 56 | |
| | | | × | Þ | ; | 00.602-1.6.111 | |
| | | | 4 | < | * | N1=40, N2=31 | × |
| | | | | | | Fold Change: 2.24 | |
| 440 | 832 | AT072873 | A27 07+1 212 5A | | | r-vame; .0104;/ | |
| | | | 117 661 1010 | | 436.16+/-215 | 436.16+/-215 | 436.16+/-215 |
| | | | 11/.304/-101.2 | | 164.58+/-155.74 | 126.68+/-109.81 | 75.01+/-128.01 |
| | | | NI=40, NZ=168 | × | N1=40, N2=6 | N1=40, N2=31 | N1=40 N2=10 |
| | | | Fold Change: 4.55 | | Fold Change: 3.37 | Fold Change: 4.14 | Dold Changes 9 70 |
| 1 | | | P-value: 0 | | P-value: 021 | D-malue: 0 | Dard Change: 6.76 |
| 441 | 836 | AI979261 | - | | | 145 88+/-125 11 | r-value: .00000 |
| | | | | | | 11:071-1:00:00 | |
| | | | × | × | ۶ | M1-40 NP-01 | ; |
| | | | | ! | \$ | 10=7N, 04-1N | × |
| | | | | | | Fold Change: 2.17 | |
| | | | | | | P-value: .00002 | |

| ie III | F. 8 | ٩ | 2.2 | 52 | 80. | 32 | 10 | 7.49 | | 11 | 66 | ٩ | 2.93 | | | | | | | | | | | | 89 | 66 | 10 | 3.01 | 002 | .07 | .55 | 9 | 2.12 |
|---------------------|------------------------------------|--------------|-------------------|-----------------|-----------------|----------------|---------------|-------------------|------------|----------------|----------------|--------------|-------------------|------------|-----------------|----------------|-------------|-------------------|-----------------|---------------|-----------------|--------------|------------------|-----------------|-----------------|----------------|---------------|-------------------|-----------------|-----------------|-----------------|--------------|-------------------|
| Normal vs Stage III | 352.73+/-154.77 839.63+/-451.94 | N1=39, N2=6 | Fold Change: 2.2 | P-value: .00452 | 281.02+/-338.08 | -25.74+/-23.32 | N1=40, N2=10 | Fold Change: 7.49 | P-value: 0 | 243.4+/-112.77 | 81.33+/-38.99 | N1=39, N2=6 | Fold Change: 2.93 | P-value: 0 | | | × | | | | | × | | | 446.58+/-157.69 | 148.67+/-68.99 | N1=40, N2=10 | Fold Change: 3.01 | P-value: .00002 | 576.75+/-132.07 | 296.15+/-132.55 | N1=40, N2=10 | Fold Change: 2.12 |
| Normal vs Stage II | 352,73+/-154,77 | N1=39, N2=31 | Fold Change: 2.11 | P-value: 0 | 281.02+/-338.08 | -12,28+/-33,2 | N1=40, N2=31 | Fold Change: 6.91 | P-value: 0 | 243.4+/-112.77 | 111.29+/-55.01 | N1=39, N2=31 | Fold Change: 2.12 | P-value: 0 | | | × | | | 35.21+/-34.58 | 207.82+/-290.06 | N1=40, N2=31 | Fold Change: 2.5 | P-value: .00091 | 446.58+/-157.69 | 182.45+/-74.24 | N1=40, N2=31 | Fold Change: 2.41 | P-vaine: 0 | | | × | |
| Normal vs Stage I | | × | | | 281.02+/-338.08 | 6.46+/-26.18 | N1=40, N2=6 | Fold Change: 6.38 | P-value: 0 | | | × | | | 160.57+/-119.71 | 240.25+/-/8.81 | N1=40, N2=6 | Fold Change: 2.01 | P-value: .00445 | | | × | | | | | × | | | | | × | |
| Normal vs Malignant | | × | | | | | × | | | | | × | | | | | × | | | | | × | | | | | × | | | | | × | |
| Normal vs All | | × | | | 282.14+/-333.79 | -2.89+/-61.5 | N1=40, N2=168 | Fold Change: 6.23 | P-value: 0 | | | × | | | | | × | | | | | × | | | 446.58+/-157.69 | 203,55+/-90.87 | N1=40, N2=168 | Fold Change: 2.21 | P-value: 0 | | | × | |
| Genbank | AI982669 | | | | AI983045 | | | | | AI985653 | | | | | AI989588 | | | | | AI990483 | | | | | AL031846 | | | | | AL037368 | | | |
| Seq ID | 837 | | | | 838 | | | | | 840 | | | | | 841 | | | | | 846 | | | | | 853 | | | | | 855 | | | |
| # | 442 | | | | 443 | ! | | | | 444 | | | | | 445 | | | | | 446 | | | | | 447 | | | | | 448 | | | |

| 7 | 6 | | | | - | | |
|----------|--------|------------|-------------------|---------------------|-------------------|--------------------|---------------------|
| * | Sed ID | Cenbank | Normal vs All | Normal vs Malignant | Normal vs Stage I | Normal vs Stage II | Normal vs Stage III |
| 44 | 827 | AL037805 | 624.14+/-319.31 | | | 614.2+/-317.15 | 614.2+/-317.15 |
| | | | 281.94+/-168.3 | | | 258.58+/-142.99 | 168.59+/-62.09 |
| | | | N1=40, N2=168 | × | × | N1=40, N2=31 | N1=40, N2=10 |
| | | | Fold Change: 2.29 | | | Fold Change: 2.38 | Fold Change: 3.3 |
| 927 | | | P-value: 0 | | | P-value: 0 | P-value: 0 |
| Ę. | 200 | ALU39445 | | | | | 103.65+/-39.97 |
| | | | ; | , | | | 204.36+/-54.58 |
| | | | < | × | × | × | N1=40, N2=10 |
| | | | | | | | Fold Change: 2.03 |
| 127 | 924 | 4 7 020000 | 40 000 000 | | | | P-value: 0 |
| 121 | 100 | ALU398/0 | 229.33+/-119.68 | | | 226.4+/-119.79 | 226.4+/-119.79 |
| | | | 104.08+/-62.24 | i | | 95.07+/-69.54 | 85.61+/-45.98 |
| | | | N1=40, N2=168 | × | × | N1=40, N2=31 | N1=40, N2=10 |
| | | | Fold Change: 2.19 | | | Fold Change: 2.41 | Fold Change: 2.58 |
| 450 | 630 | AT 020017 | r-value: 0 | | | P-value: 0 | P-value: .0005 |
| 70 | 700 | AL05991/ | | | | | 190.41+/-139.22 |
| • | | | Þ | ; | | | 427.64+/-204.38 |
| | | | × | × | × | × | N1=40, N2=10 |
| | | | | | | | Fold Change: 2.48 |
| 1 | ,,,, | | | | | | P-value: .00467 |
| <u>ş</u> | 804 | AL040178 | 277.18+/-128.12 | | 277.18+/-128.12 | 277.18+/-128.12 | 277.18+/-128.12 |
| | | | /3.94+/-56.64 | | 106.38+/-90.67 | 59.11+/-47.84 | 44.44+/-29.67 |
| | | | N1=39, N2=168 | × | N1=39, N2=10 | N1=39, N2=31 | N1=39, N2=6 |
| | | | Fold Change: 3.48 | | Fold Change: 2.79 | Fold Change: 3.76 | Fold Change: 4.57 |
| 454 | 270 | 47 040040 | r-value: 0 | | P-value: .01209 | P-value: 0 | P-value: 0 |
| 10 | 202 | AL040912 | 511.14+/-137.52 | | | 304.56+/-132.78 | 304.56+/-132.78 |
| | | | 80,94+/-86.46 | i | | 69.8+/-53.68 | 52.07+/-61.09 |
| | | | NI=40, NZ=168 | × | × | N1=40, N2=31 | N1=40, N2=10 |
| | | | Fold Change: 4.08 | | | Fold Change: 4.66 | Fold Change: 6.69 |
| | | | P-value: 0 | | | P-value: 0 | P-value: .00001 |
| 52 | 866 | AL041815 | 257.28+/-104.74 | | | 257.35+/-106.11 | |
| | | | 129.57+/-68.69 | ; | | 124.69+/-56.27 | |
| | | | NI=40, NZ=168 | × | × | N1=40, N2=31 | × |
| | | | Fold Change: 2.04 | | | Fold Change: 2 | |
| | | | 1-vaine. 0 | | | P-value: 0 | |

| Seq ID | Genbank | Normal vs All | Normal vs Malignant | Normal vs Stage I | Normal vs Stage II | Normal vs Stage III |
|--------|----------|--------------------|---------------------|-------------------|--------------------|---------------------|
| | AL042492 | 801.96+/-843.5 | | 809.69+/-853.09 | 809.69+/-853.09 | 806.69+/-853.09 |
| | | 56.95+/-101.91 | | 101.65+/-170.57 | 40.59+/-106.87 | 11.79+/-18.8 |
| | | N1=40, N2=168 | × | N1=40, N2=6 | N1=40, N2=31 | N1=40, N2=10 |
| | | Fold Change: 12.71 | | Fold Change: 8.78 | Fold Change: 15 | Fold Change: 20.99 |
| - | | P-value: 0 | | P-value: .00362 | P-vafue: 0 | P-value: 0 |
| | AL042923 | | | | | 3125.4+/-1239.9 |
| | | , | į | 1 | | 1575.01+/-724.4 |
| | | × | × | × | × | N1=40, N2=10 |
| | | | | | | Fold Change: 2.07 |
| ľ | AL043980 | | | | 498.96+/-198.25 | 498.96+/-198.25 |
| | | | | | 248.27+/-101.89 | 206.45+/-96.76 |
| | | × | × | × | N1=40, N2=31 | N1=40, N2=10 |
| | | | | | Fold Change: 2.06 | Fold Change: 2.43 |
| | 2000 | | | | P-value: 0 | P-value: .00002 |
| • | AL044366 | 137,98+/-186,9 | | | | |
| | | N1=40 N2=168 | × | ۲ | > | Þ |
| | | Fold Change: 2.23 | • | 4 | < | < |
| | | P-value: .00018 | | | | |
| * | AL044613 | | | | | 304+/-112.89 |
| | | ; | 1 | | | 102.49+/-79.71 |
| | | × | × | × | × | N1=40, N2=10 |
| | | | | | | Fold Change: 3.15 |
| ľ | | | | | | P-value: .00076 |
| ~ | AL046941 | 425.75+/-236.48 | | | 428.58+/-238.89 | 428.58+/-238.89 |
| | | 53.79+/-102.7 | | | 34.11+/-80,54 | -23.64+/-33.24 |
| | | N1=40, N2=168 | × | × | N1=40, N2=31 | NI=40, N2=10 |
| | | Fold Change: 8.01 | | | Fold Change: 9.66 | Fold Change: 16.81 |
| | | P-value: 0 | | | P-value: 0 | P-value: 0 |
| • | AL046946 | | | | | 713.56+/-217.52 |
| | | ; | i | | | 287.87+/-116 |
| | | × | × | × | × | N1=40, N2=10 |
| | | | | | | Fold Change: 2.57 |
| 1 | | | | | | P-value: .00006 |

| | Genbank | Normal vs All | Normal vs Malignant | Normal vs Stage I | Normal vs Stage II | Normal vs Stage III |
|-----|----------|-------------------|---------------------|-------------------|--------------------|---------------------|
| 878 | AL048304 | | | | | 308.79+/-74.62 |
| | | × | × | × | × | N1=40, N2=10 |
| | | | | | | Fold Change: 3.99 |
| | | | | | | P-value: .00402 |
| 879 | AL048386 | 233.2+/-107.71 | | | 232.52+/-109.03 | 232.52+/-109.03 |
| | | 118.45+/-83.53 | | | 94.13+/-59.17 | 73.94+/-51.55 |
| | | N1=40, N2=168 | × | × | N1=40, N2=31 | N1=40, N2=10 |
| | | Fold Change: 2.19 | | | Fold Change: 2.63 | Fold Change: 3.32 |
| | | P-value: 0 | | | P-value: 0 | P-value: ,00019 |
| 880 | AL048399 | | | | 768,94+/-280,35 | 768.94+/-280.35 |
| | | | | | 336,8+/-159,66 | 378.81+/-201.68 |
| | | × | × | × | N1=40, N2=31 | N1=40, N2=10 |
| | | | | | Fold Change: 2.37 | Fold Change: 2.22 |
| | | | | | P-value: 0 | P-value: .00292 |
| 881 | AL048962 | 951.97+/-353.33 | | | 944+/-354.29 | 944+/-354.29 |
| | | 498.61+/-346.17 | | | 469.14+/-334.1 | 400.6+/-283.17 |
| | | N1=40, N2=168 | × | × | N1=40, N2=31 | N1=40, N2=10 |
| | | Fold Change: 2.03 | | | Fold Change: 2.17 | Fold Change: 2,45 |
| | | P-value: 0 | | | P-value: 0 | P-value: .00012 |
| 883 | AL049257 | | | | 258.77+/-104.32 | 258.77+/-104.32 |
| | | | | | 110,97+/-53.29 | 104.27+/-37.82 |
| | | × | × | × | N1=40, N2=31 | N1=40, N2=10 |
| | | | | | Fold Change: 2.35 | Fold Change: 2.34 |
| | | | | | P-value: 0 | P-value: 0 |
| 884 | AL049423 | | | | 385.45+/-146.45 | 385.45+/-146.45 |
| | | | | | 176.43+/-82.21 | 131.88+/-33.79 |
| | | × | × | × | N1=40, N2=31 | N1=40, N2=10 |
| | | | | | Fold Change: 2.19 | Fold Change: 2.73 |
| | | | | | P-value: 0 | P-value: 0 |
| 882 | AL049471 | | | | 585.55+/-164.83 | 585.55+/-164.83 |
| | | | | | 312.66+/-134.95 | 304.04+/-127.09 |
| | | × | × | × | N1=40, N2=31 | N1=40, N2=10 |
| | | | | | Fold Change: 2.01 | Fold Change: 2.01 |
| | | | | | P-value: 0 | P-vrahre: 00064 |

| September Normal vs Stage | # | Sed III | Genbank | Normal vs All | Mouracol we Mark | | | |
|--|-----|---------|-----------|-------------------|--------------------|-------------------|--------------------|----------------------------|
| 888 ALA69957 X X X 218.014-118.84 889 ALA69957 X X X X 18.014-118.84 890 ALA69067 X X X X X X X 18.04-116.51 890 ALA69067 257.594-77.75 894 ALA79779 131.404-126.87 894 ALA79779 131.404-128.97 896 ALA79779 131.404-128.97 896 ALA79779 131.404-128.97 897 ALA79779 138.84-138.91 897 ALA79779 198.84-138.91 898 ALA79780 198.84-138.91 899 ALA79780 198.84-138.91 890 ALA79780 198.84-138.91 890 ALA79780 198.84-138.91 891 ALA79780 198.84-138.91 892 ALA79780 198.84-138.91 893 ALA79780 198.84-138.91 894 ALA79780 198.84-138.91 895 ALA79780 198.84-138.91 896 ALA79780 198.84-138.91 897 ALA79780 198.84-138.91 898 ALA79780 198.84-138.91 899 ALA79780 198.84-138.91 890 ALA79780 198.84-138.91 | 470 | ľ | AL049949 | TO SE TRIVES | Normal vs Mangnant | Normal vs Stage I | Normal vs Stage II | Normal vs Stage III |
| See | | | | | | | 525.51+/-234.08 | 525.51+/-234.08 |
| 888 AIA61957 | | | | ۶ | i | | 218.01+/-118.84 | 184.51+/-151.36 |
| See Al. Al. Al. 9597 See Al. Al. 9797 See Al. 9797 | | | | < | × | × | N1=40, N2=31 | N1=40 N2=10 |
| See | | | | | | | Fold Change: 2.27 | Fold Change: 3.21 |
| September Sept | E | 888 | AL049957 | | | | P-value: 0 | P-value: .00178 |
| National Changes 1.29 National Changes 1.29 | | | | | | | 664.06+/-250.51 | |
| Solution | | | | > | ; | | 339.26+/-164.3 | |
| See | | | | 4 | * | × | NI=39, N2=31 | × |
| Sept AL693062 | | | | | | | Fold Change: 1.98 | ; |
| X | 12 | 890 | AT.050002 | | | | P-value: 0 | |
| Section | | ı İ | - | | | | | 320 05+/-114 06 |
| National Character Nationa | | | | , | | | | 152 33±/ 60 11 |
| B93 AL689367 227/594-77/75 1554-47068 X | | | | < | × | × | × | M1=40 N2-10 |
| 157-1594-7735 157-1594-773 | | | | | | | ! | D-11 C |
| 894 AL07970 5157594-7773 27594-7775 894 AL07970 5164-120.63 X X 117464-7636 894 AL07970 5164-139.76 315494-139.76 31644-7536 895 AL07970 7584-43.88 1144-76.89 81144-76.89 867.794-51.4 896 AL07970 7584-43.89 X 1044-76.89 867.794-51.4 897 AL07970 182-24-25.80 X 1044-72.80 867.794-51.4 897 AL07970 182-24-25.80 X 1044-72.80 87.794-22.7 898 AL07970 182-24-25.80 X 1044-72.80 87.794-22.7 899 AL07970 182-24-25.80 X 1044-72.80 87.794-22.7 897 AL07970 182-24-25.80 X 1044-72.80 87.794-22.7 899 AL07970 182-24-25.80 X 1044-72.80 87.794-22.7 897 AL07970 182-24-25.80 X 1044-72.80 87.794-22.7 897 AL07970 182-24-25.80 X 1044-72.80 87.794-22.7 898 AL07970 182-24-25.80 X 1044-72.80 87.794-22.7 899 AL07970 182-24-25.80 X 1044-72.80 87.794-22.7 899 AL07970 182-24-25.80 X 1044-72.80 87.794-22.7 890 AL07970 182-24-25.80 X 1044-72.80 87.794-22.7 890 AL07970 182-24-25.80 X 1044-72.80 87.794-22.7 890 AL07970 182-24-25.80 X 1044-72.80 87.794-22.7 890 AL07970 182-24-25.80 X 1044-72.80 87.794-22.7 890 AL07970 182-24-25.80 X 1044-72.80 87.794-22.7 890 AL07970 182-24-25.80 X 1044-72.80 87.794-22.7 890 AL07970 182-24-25.80 X 1044-72.80 87.74-22.80 87 | ١, | 000 | | | | | | Fold Change; 2.08 |
| 1554-150.63 | , | 673 | AL050367 | 257.59+/-77.75 | | | 200 000 1:00 000 | r-vaine: .0001/ |
| Ni Al Ni Ni Al Al Ni Al Ni Al Ni Al Al Al Al Al Al Al A | | | | 155+/-120,63 | | | 411 40: (200 | |
| Fold Changer 2 | | | | N1=40, N2=168 | ٢ | Þ | 11/.104/-/0.86 | |
| Profiles 0.1 Profiles 0.2 Profiles 0.4 10.2024-6.58 13.494-189.76 13.494-189.76 13.494-189.76 10.2024-6.58 13.494-189.76 13.494-189.76 13.494-189.76 10.2024-6.58 13.494-189.76 13.494-189.76 13.494-189.76 10.2024-6.58 13.494-189.76 13.494-189.76 13.494-189.76 10.2024-6.58 13.494-189.76 13.494-189.76 13.494-189.76 10.2024-6.58 13.494-18.76 13.494-18.76 13.494-18.76 10.2024-6.58 13.494-18.76 13.494-18.76 13.494-18.76 10.2024-6.58 13.494-18.76 13.494-18.76 13.494-18.76 10.2024-6.58 13.494-18.76 13.494-18.76 13.494-18.76 10.2024-6.2024 13.494-18.76 13.494-18.76 10.2024-6.2024 13.494-18.76 13.494-18.76 10.2024-6.2024 13.494-18.76 13.494-18.76 10.2024-6.2024 13.494-18.76 13.494-18.76 10.2024-6.2024 13.494-18.76 13.494-18.76 10.2024-6.2024 13.494-18.76 13.494-18.76 10.2024-6.2024 13.494-18.76 13.494-18.76 10.2024-6.2024 13.494-18.76 13.494-18.76 10.2024-6.2024 13.494-18.76 13.494-18.76 10.2024-6.2024 13.494-18.76 13.494-18.76 10.2024-6.2024 13.494-18.76 13.494-18.76 10.2024-6.2024 13.494-18.76 13.494-18.76 13.494-18.76 10.2024-6.2024 13.494-18.76 13.494-18.76 10.2024-6.2024 13.494-18.76 13.494-18.76 10.2024-6.2024 13.494-18.76 13.494-18.76 10.2024-6.2024 13.494-18.76 13.494-18.76 10.2024-6.2024 13.494-18.76 13.494-18.76 10.2024-6.2024 13.494-18.76 13.494-18.76 10.2024-6.2024 13.494-18.76 13.494-18.76 10.2024-6.2024 13.494-18.76 13.494-18.76 13.494-18.76 10.2024-6.2024 13.494-18.76 13.494-18.76 13.494-18.76 10.2024-6.2024 13.494-18.76 13.494-18.76 13.494-18.76 10.2024-6.2024 13.494-18.76 13.494-18.76 13.494-18.76 10.2024-6.2024 13.494-18.76 13.494-18.76 13.494-18.76 10.2024-6.2024 13.494-18.76 13.494-18.76 13.494-18.76 13.494-18.76 13.494-18.76 13.494-18.76 13.494-18.76 13.494-18.76 13.494-18.76 13.494-1 | | | | Fold Change: 2 | \$ | * | N1=40, N2=31 | × |
| 894 ALITY279 313.494+189.76 313.494+139.76 313.494+139.76 313.494+139.76 313.494+139.76 313.494+139.76 313.494+139.76 313.494+139.76 313.494+139.76 313.494+38.87 313.494+38.77 | - 1 | | | P-value: 0 | | | Fold Change: 2.49 | |
| 108.294-85.88 1344-1487.06 1343-444-1887.06 1344-1487.06 1343-444-1887.06 1344-148 | | 894 | AT.079279 | 313 AOL/ 100 76 | | | P-value: 0 | |
| No. 2017 | | | | 100 101 101 00 | | 313.49+/-189.76 | 313.49+/-189.76 | 313 49+/-180 76 |
| Number N | | | | MI 40 3 m 160 | | 114+/-76.08 | 86.73+/-51.4 | 49 04+/-28 4 |
| Post Change 2.87 Pod Change 2.87 Pod Change 2.80 | | | | D-13 CT 0.00 | × | N1=40, N2=6 | NI=40, N2=31 | NI=do NO=10 |
| 896 ALOPYOT 2-8014-7224-3 2-100-14-202-3 2-10-14-20-4 | | | | roid Change: 2.87 | | Fold Change: 2.51 | Fold Change: 3.29 | Rold Change: 5 60 |
| Control Cont | 1, | 896 | AT 070707 | r-vame: 0 | | P-value: .00823 | P-value: 0 | P-value: 0 |
| No. 2007 | | | 1016107 | 75.97.1-7.24.37 | | 261.69+/-226.08 | 261.69+/-226.08 | 261 60±/,226 08 |
| Ni = 0, Vi > 0 | | | | 07.8047-48.3 | | 82.89+/-41.99 | 48.17+/-30.27 | 70 831/ 71 76 |
| Fold Change: 3.55 Fold Change: 2.13 Fold Change: 2.14 | | | | N1=40, N2=168 | × | N1=40, N2=6 | N1=40 N2=31 | 02.12-40.81 MI-40.80 30 |
| 897 AL079769 19-20-20-20-20-20-20-20-20-20-20-20-20-20- | | | | Fold Change: 3.55 | | Fold Change: 2 73 | Fold Change: 4 62 | 14-40, IAZ-10 |
| 897 AL079769 1198_264-1265.48 2013-44-2656.02 2013-44-266.02 2013 | | | | P-value: 0 | | P-value: 0013 | Parling 4.03 | Fold Change: 2.7 |
| X X X X X X X X X X X X X X X X X X X | | 897 | AL079769 | 198.26+/-265.48 | | - tajuo0013 | r-value: 0 | P-value: 0 |
| X 451,540-269.91 X N1-40, N2-31 Fold Change, 2.73 | | | | 337.38+/-218.01 | | | 201./4+/-268.02 | 201.74+/-268.02 |
| A N1=40, N2=31 Fold Change: 2.73 | | | | N1=40, N2=168 | ۶ | ; | 451.36+/-269.91 | 415.85+/-142.87 |
| Fold Change: 2.73 | | | | Fold Change: 2 06 | • | * | N1=40, N2=31 | N1-40, N2-10 |
| | | | | P-value 0 | | | Fold Change: 2.73 | Fold Change: 2.88 |

| 889 AL079949 X X X X X INTEGERAL STATE OF THE PROBLEM STATE OF THE PROBL | 889 AL079949 X X X X X X X X X X X X X | # | Seq ID | Genbank | Normal vs All | Normal vs Malignant | Normal ve Stage I | Normal va Ctage II | Manney |
|--|--|-----|--------|------------|-------------------|---------------------|---------------------|---------------------|---------------------|
| September Action | State Stat | 477 | 868 | AI.079949 | | 4 | A CONTROL TO CHARGO | MOTHER VS STABLE II | Normal vs Stage III |
| State | Sign ALABB192 | | } | | | | | 382.36+/-135.98 | 382,36+/-135,98 |
| Sign ALABBI 2 | Solid Changer 221 Not Changer 221 Not Changer 221 Not Changer 221 Not Changer 221 Not Changer 221 Not Changer 221 Not Changer 221 Not Changer 221 Not Changer 222 Not Changer 224 Not Changer 225 Not Chan | | | | ì | | | 172.14+/-65.04 | 160.74+/-65.7 |
| Separate | Signature Sign | | | | × | × | × | N1=40, N2=31 | NI=40. N2=10 |
| 899 AL080192 | 899 AL080192 X | | | | | | | Fold Change: 2.21 | Fold Change: 2.34 |
| 10.24 1.02 | National State | 2 | 000 | - T 000100 | | | | P-value: 0 | P-value: .00001 |
| March Marc | March Marc | 0 | 660 | 76T020TW | | | | 103.54+/-69.27 | |
| March Marc | March Marc | | | | , | | | 242.17+/-111.66 | |
| March Marc | Section | | | | × | × | × | N1=40, N2=31 | × |
| 903 Ali20446 304,544-95.79 N-value: 0 | 903 Ali20446 304,544-95.79 N-value: 0 | | | | | | | Fold Change: 2.54 | |
| 914 AW00350 150.454-05.79 | 914 AW00389 255.54-12.34 | 1 | | | | | | P-value: 0 | |
| 151.0F4-0.75 151. | 151.0F4-51.13 | 2 | 302 | AL120446 | 304.54+/-95.79 | | | 302.81+/-96.4 | 302.81+/-96.4 |
| New Color | NW00580 NW00 | | | | 151.014/-51.13 | | | 143,68+/-46,25 | 137.13+/-46.57 |
| Fold Change; 2 Fold | Fold Changer 2 Fold | | | | N1=40, N2=168 | × | × | NI=40. N2=31 | N1=40 N2=10 |
| 994 AW000899 15-74-12-34 AV000899 15-74-12-34 AV000899 15-74-12-34 AV000899 15-74-12-34 AV000899 15-74-12-34 AV000899 15-74-12-34 AV000896 17-74-12-34 AV000896 17-74-12-34 AV000896 18-74-12-34 AV0008918 X X X X X X X X X X X X X X X X X X X | Project Proj | | | | Fold Change: 2 | | | Hold Changes, 2 07 | 01 211 01 11 |
| 904 AW000899 255.574-123.44 25.254-1123.44 25.2 | 904 AW000899 255.53+1123.44 10.23.65+1173.34 10.23.65+1173.34 10.25.65+1173.34 10.25.65+1173.44 10.73.65+1173.44 1 | | | | P-value: 0 | | | Four Change: 2.0/ | rold Change: 2.16 |
| 12.545+1073 12.545+1074 12.545+1075 12.545+1075 12.545+1075 13.545+10.244 13.545 | 112.544/1073 X X X X X X X X X | 8 | 904 | AW000899 | 255.52+/-122.34 | | | Der Coll 100 0 | r-vaue: .00001 |
| Ni - 20, N | Ni - 25, Vi > 10, V | | | | 125.26+/-107.3 | | | 255.52+/-122.34 | |
| Fold Changer 2.23 | Field Change 2.23 Field Change 2.24 | | | | N1=39, N2=168 | þ | Þ | 130.124-122.05 | • |
| March Marc | Production | | | | Rold Changes 2 22 | < | 4 | NI=39, NZ=31 | × |
| 907 AW002846 222.394-i159506 March 1554 | 907 AW00246 122-24-1159 of Prepare 00003 907 AW00246 X X X X N1-40, N2-468 X X X N1-40, N2-468 X X X N1-40, N2-468 X X X N1-40, N2-468 X X N1-40, N2-468 X X N1-40, N2-46 X X N1-40, N2-47 X X N1-40, N2-47 X X N1-40, N2-47 X X N1-40, N2-47 X X N1-40, N2-47 X X X N1-40, N2-47 X N1-40, N2-47 X X N1-40, N2-57 X N1-40, N2-57 X N1-40, N2-57 X N1-40, N2-57 X X X N1-40, N2-57 X N1-40, N2-57 X N1-40, N2-57 X X N1-40, N2-57 X N1-40, N2-57 X X X X X X X X X X X X X X X X X X X | | | | Pavalue: 0 | | | Fold Change: 2.06 | |
| AV00540 AX AV005418 AV005 | Available Avai | 2 | 200 | ANDOROGA | 200 201 (200 00 | | | P-value: .00003 | |
| 159.277+1234 | 100 | 5 | Ř | AW002046 | 282.39+/-199.06 | | | | 283.14+/-201.6 |
| Nation N | Number N | | | | 139.23+/-115.04 | | | | 96.89-/+8.76 |
| 912 AW003562 P-ralbac 0 211579-48124 X X N1=94, N2=90 X Pold Change: 2.23 Pold Change: 2.23 Pold Change: 2.24 AW00518 X X X X X X X X X X X X X X X X X X X | 912 AW003462 P-rulbine 0 211.574-481.24 | | | | NI=40, NZ=168 | × | × | × | N1=40 N2=10 |
| 912 AW003362 F-value: 0 211.574-531.24 X X N1=59, N2=10 Rod Change: 2.22 Fold Change: 2.22 X X N4=6, N2=10 Fold Change: 2.22 X X X N4=6, N2=10 Fold Change: 2.24 X X X X N4=6, N2=11 Fold Change: 2.41 Fold Change: 2.41 Fold Change: 2.41 | 912 AW003362 F-values 0 211.574-81.24 X X 89.064-83.95 Y X N 1-39, N2-40 Pold Change: 2.22 P-value: 0.2046 205.574-7.234.3 X X X N 1-40, N2-51 Fold Change: 2.21 Fold Change: 2. | | | | Fold Change: 2.15 | | | | Fold Change: 2.7 |
| 912 AW003562 X 2115745124 Size State | 912 AW003562 X 211574-8124 X X 8064-5355 X N1-59, N2-10 Fold Change: 2.22 914 AW005418 X X X X X X X X X X X X X X X X X X X | 1 | | | P-value: 0 | | | | P-value: 00045 |
| X | No. | 82 | 912 | AW003362 | | | 211.57+/-81.24 | - | 2000 |
| N N N N N N N N N N | N N N N N N N N N N | | | | ; | | 89.06+/-83.95 | | |
| Pold Chapper 2.22 Pold Chapper 2.22 Pold Chapper 2.22 Pold Chapper 2.22 Pold Chapper 2.22 Pold Chapper 2.23 N | 914 AW005418 Fold Changes 2.22 Powlate, 02046 205.574/234.3 X X X X X X N 1-40, 102-51 Fold Changes, 2.41 Fo | | | | × | × | N1=39, N2=10 | × | × |
| 914 AW005418 Pevalue: 02046 205.574/234.3 (6.114-110.71 X X X X NI-04.110.71 Pold Change, 2.41 Pold Change, 2.41 | 914 AW005418 P-value: 02046 205.57+/234.3 X X X X X X X X X X A A | | | | | | Fold Change: 2.32 | | ; |
| 914 AW005418 205.574/234.3 | 914 AW005418 205.574-7234.3 917 AW005418 705.774-7234.3 87 N1-46,10,71 Pold Change, 2,41 Pold Change | l | | | | | P-value: .02046 | | |
| X | X X (3.14-110.71 N1-40, N2-51 Fold Canger, 2.41 F-value; 0.0083 | ç | 914 | AW005418 | | | | 205.57+/-234.3 | 205.57+/-234.3 |
| X N1=40, N2=31 Fold Change: 2.41 | X X N1=40, N2=51 Pold Change; 2,41 Polds: ,00083 P-value; ,00083 | | | | ÷ | ; | | 67.1+/-110.71 | 10.33+/-83.48 |
| | | | | | < | × | × | N1=40, N2=31 | N1=40, N2=10 |
| | | | | | | | | Fold Change: 2.41 | Fold Change: 3.7 |

| ĺ | | | | | | | |
|----------------|--------|----------|------------------------------------|---------------------|------------------------------------|--------------------|---------------------|
| # | Seq ID | Genbank | Normal vs All | Normal vs Malignant | Normal vs Stage I | Normal vs Stage II | Normal vs Stage III |
| 484 | 915 | AW005814 | | | 146.08+/-90.43 | C | |
| | | | × | × | N1=40, N2=6 | × | × |
| | | | | | Fold Change: 2.4 P-value: 00072 | | |
| 485 | 916 | AW006235 | 344.79+/-207.97 | | 346,9+/-210.26 | 346,9+/-210.26 | 346.9+/-210.26 |
| | | | 103.81+/-60.65 | | 126.57+/-35.39 | 95.72+/-56.93 | 99.92+/-23.08 |
| | | | N1=40, N2=168 | × | N1=40, N2=6 | N1=40, N2=31 | N1=40, N2=10 |
| | | | Fold Change: 3.17 | | Fold Change: 2.35 | Fold Change: 3.43 | Fold Change: 2,94 |
| | - 1 | | P-value: 0 | | P-value: .00014 | P-value: 0 | P-value: 0 |
| 486 | 919 | AW006898 | 841.88+/-394.55 | | | 835.42+/-397.56 | 835.42+/-397.56 |
| | | | 331.12+/-203.65 | | | 305.25+/-163.97 | 305.65+/-212.32 |
| | | | N1=40, N2=168 | × | × | N1=40, N2=31 | N1=40, N2=10 |
| | | | Fold Change: 2.68 | | | Fold Change: 2.7 | Fold Change: 2.91 |
| 1 | - 1 | | F-vame: 0 | | | P-value: 0 | P-value: .00883 |
| 487 | 920 | AW006998 | | | | 79.75+/-104.09 | |
| | | | | | | 221.43+/-283.12 | |
| | | | × | × | × | N1=40, N2=31 | × |
| | | | | | | Fold Change: 2.76 | |
| | | | | | | P-value: .00002 | |
| 488 | 921 | AW007080 | 226.13+/-116.85 | | | 223.2+/-116.87 | 223,2+/-116,87 |
| | | | 62.58+/-55.91 | | | 55.48+/-39.27 | 48.27+/-49.28 |
| | | | N1=40, N2=168 | × | × | N1=40, N2=31 | N1=40, N2=10 |
| | | | Fold Change: 3.65 | | | Fold Change: 3.79 | Fold Change: 4.6 |
| | | | F-value: 0 | | | P-value: 0 | P-value: .00008 |
| 8 6 | 925 | AW007586 | 552.57+/-493.88 992.94+/-672.01 | | | 552.57+/-493.88 | |
| | | | N1=40, N2=168 | × | × | N1=40. N2=31 | × |
| | | | Fold Change: 2.04 | | | Fold Change: 2.99 | |
| | - 1 | | P-value: .00003 | | | P-value: 0 | |
| 6 | 726 | AW007983 | 308.57+/-180.09 | | 308.57+/-180.09 | 308.57+/-180.09 | 308.57+/-180.09 |
| | | | 124.89+/-104.9 | | 120.02+/-133.67 | 98.43+/-60.3 | 57.05+/-49.19 |
| | | | N1=39, N2=168 | × | N1=39, N2=10 | NI=39, N2=31 | N1=39, N2=6 |
| | | | Fold Change: 2.45 | | Fold Change: 3.11 | Fold Change: 2.82 | Fold Change: 4.11 |
| | | | - Anino | | r-vailleU2042 | r-value: 0 | F-vaine: 0 |

| # | Seq ID | Genbank | Normal vs All | Normal vs Malignant | Normal vs Stage I | Normal vs Stage II | Normal vs Stage III |
|-----|--------|----------|-------------------|---------------------|--------------------------------------|--------------------|-------------------------------------|
| 491 | 929 | AW009505 | | | | | 210.48+/-63.01 |
| | | | × | × | × | × | N1=40, N2=10 |
| | | | | | | | Fold Change: 2.06 P-value: 00023 |
| 492 | 930 | AW013949 | | | 228.98+/-140.11 | | |
| | | | × | × | N1=40, N2=6 | × | × |
| | | | | | Fold Change: 2.49 P-value: .00743 | | |
| 493 | 932 | AW014647 | 222.93+/-95.23 | | | 222.93+/-95.23 | 222,93+/-95.23 |
| | | | 93.53+/-66.06 | | | 87.53+/-67.66 | 103.61+/-77.88 |
| | | | N1=39, N2=168 | × | × | N1=39, N2=31 | N1=39, N2=6 |
| | | | Fold Change: 2.35 | | | Fold Change: 2.48 | Fold Change: 2.1 |
| | | | P-value: 0 | | | P-value: 0 | P-value: ,00171 |
| 464 | 933 | AW014764 | 299.88+/-160.3 | | | 299.88+/-160.3 | |
| | | | 148.29+/-123.67 | | | 143.2+/-104.38 | |
| | - | | N1=39, N2=168 | × | × | N1=39, N2=31 | × |
| | | | Fold Change: 2.02 | | | Fold Change: 2.02 | |
| _ | | | P-value: 0 | | | P-vame: 0 | |
| 495 | 934 | AW015571 | | | | | 648,86+/-400,58 |
| | | | | | | | 174.43+/-228.74 |
| | | | × | × | × | × | N1=40, N2=10 |
| | | | | | | | Fold Change: 6.15 |
| 496 | 938 | AW021108 | 220 45+/-143.17 | | | 220.45+/-143.17 | 220.45+/-143.17 |
| : | | | 97.68+/-75.55 | | | 87.82+/-65,1 | 76.42+/-41.74 |
| | | | N1=39, N2=168 | × | × | N1=39, N2=31 | N1=39, N2=6 |
| | | | Fold Change: 2.37 | | | Fold Change: 2.67 | Fold Change: 2.68 |
| | | | P-value: 0 | | | P-value: 0 | P-value: .00045 |
| 497 | 939 | AW021169 | 256.18+/-141.26 | | | | 254.1+/-142.48 |
| | | | 128.88+/-175.65 | | | | 131+/-172.12 |
| | | | N1=40, N2=168 | × | × | × | N1=40, N2=10 |
| | | | Fold Change: 2.2 | | | | Fold Change: 2.58 |
| | | | P-value: 0 | | | | P-value: .00997 |

| 4 | Coo ID | Conhonly | Mormol ve All | Normal ve Molimont | Normal ve Stone I | Normal ve Stage II | Normal we Stage III |
|-----|--------|----------|-------------------|-----------------------|-------------------|---------------------|---------------------|
| - | nace . | Colloann | LIOTATIAL VS ALL | AND ING TO INTERBURIE | MORNING 13 STUBER | TADEMAI AS SINGE TO | TOTAL TO STUBE IT |
| 498 | 941 | AW022607 | | | | 820.58+/-231.5 | 820.58+/-231.5 |
| | | | | | | 436,42+/-247.61 | 384.68+/-242.39 |
| | | | × | × | × | N1=40, N2=31 | NI=40, N2=10 |
| | | | | | | Fold Change: 2.16 | Fold Change: 2,39 |
| | | | | | | P-value: 0 | P-value: .00116 |
| 499 | 943 | AW023188 | | | | 290.36+/-132.68 | |
| | | | | | | 130,36+/-49.53 | |
| | | | × | × | × | N1=39, N2=31 | × |
| | | | | | | Fold Change: 2.07 | |
| 200 | 976 | AW024285 | | | | r-vaine. 0 | 250 28+/-98 87 |
| 3 | | | | | | | 572.73+/-295.31 |
| | | | × | × | × | × | N1=40, N2=10 |
| | | | | | | | Fold Change: 2.27 |
| | | | | | | | P-value: .00009 |
| 201 | 946 | AW024434 | 790.91+/-331.85 | | | 789.94/-336.13 | |
| | | | 432.05+/-300.27 | | | 。 365.21+/-203.57 | |
| | | | N1=40, N2=168 | × | × | N1=40, N2=31 | × |
| | | | Fold Change: 2.03 | | | Fold Change: 2.21 | |
| | | | P-value: 0 | | | P-value: 0 | |
| 22 | 948 | AW024795 | | | | | 194.61+/-162.76 |
| | | | | | | | 374.26+/-130.52 |
| | | | × | × | × | × | N1=40, N2=10 |
| | | | | | | | Fold Change: 2.22 |
| | | | | | | | P-value: .00004 |
| 503 | 325 | AW044663 | | | | | 216.92+/-78.22 |
| | | | | | | | 104,46+/-39.86 |
| | | | × | × | × | × | N1=40, N2=10 |
| | | | | | | | Fold Change: 2.01 |
| | | | | | | | P-value: .00001 |
| 504 | 953 | AW051492 | 440.5+/-328.97 | | | 442.65+/-332.99 | 442.65+/-332.99 |
| | | | 200.34+/-109.37 | , | | 174.66+/-95.13 | 133.38+/-80.06 |
| | | | N1=40, N2=168 | × | × | N1=40, N2=31 | N1=40, N2=10 |
| | | | Fold Change: 2.03 | | | Fold Change: 2.4 | Fold Change: 3.04 |
| | | | P-value: 0 | | | P-value: 0 | P-value: .00008 |

| Approximately and the second s | Normal vs Malignant Normal vs Stage I Normal vs Stage II Normal vs Stage III | 200.32+/-119.43 200.32+/-119.43 | | X N1=39, N2=10 N1=39, N2=31 | Fold Change: 2.54 Fold Change: 2.68 Fe | P-value: .0089 P-value: 0 | 470.75+/-309.62 470.75+/-309.62 | | X N1=40, N2=6 N1=40, N2=31 | Fold Change: 2.24 Fold Change: 3.13 F | P-value: .00778 P-value: 0 | | | × | • | | 231.134/-145.06 | | X N1=40, N2=31 X | Fold Change: 2.13 | F-value: .00004 | | X X | | 215 94/-67 18 | 84.24+7.31.19 | X X X | ĬŢ. | P-value: 00001 | 321.52+/-187.56 321.52+/-187.56 | 25,000,000 | - |
|--|--|---------------------------------|---------------|-----------------------------|--|---------------------------|---------------------------------|--------------|----------------------------|---------------------------------------|----------------------------|-----------------|----------------|---------------|-------------------|------------|-----------------|---|------------------|-------------------|-----------------|-----------------|---------------|---------------------------------|---------------|---------------|-------|-----|----------------|---------------------------------|------------|---|
| | Normal vs All | 3 | 68.25+/-52.36 | N1=39, N2=168 | Fold Change: 2.47 | P-value: 0 | 470.61+/-305.63 | 179.69+/-105 | N1=40, N2=168 | Fold Change: 2.63 | P-value: 0 | 233,51+/-142,32 | 116.34+/-81.47 | N1=40, N2=168 | Fold Change: 2.05 | P-value: 0 | | ł | × | | 27 027 170 | 225.06+/-178.61 | N1-40, N2-168 | Fold Change: 3.08 P-value: 0 | | | × | | | | | × |
| | a | 955 AW052186 | | | | | se C16443 | | | | | 77 C17781 | | | | | 7 D55884 | | | | D.C31777 | | | | 1 D79487 | | | | | 1 H11724 | | |
| - 1 | 1 | 505 95 | | | | - 1 | 506 956 | | | | - 1 | 507 957 | | | | | 208 967 | | | | 600 070 | | | | 510 971 | | | | | 511 981 | | |

| | | | , | | | | |
|-----|--------|---------|--|--------------------|--|--------------------|--|
| 4: | Sed ID | Genbank | Normal vs All | Normal vs Mangnant | Normal vs Stage I | Normal vs Stage II | Normal vs Stage III |
| 512 | .983 | H15868 | 347.4+/-390.17 | | 344.41+/-394.81 | | |
| | | | 933 09+71186 46 | | 967 694/1057 17 | | |
| _ | | | NI=40 N7=168 | × | A=CM ON=IM | * | > |
| | | | 11 1 1 0 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 | • | 7 11 7 7 7 7 7 7 7 7 7 7 7 7 7 7 7 7 7 | • | • |
| | | | Poid Change: 2.15 | | P-value: 02477 | | |
| 513 | 984 | H16294 | | | | | 812.95+/-387.97 |
| | | | | | | | 205 8+/-82 15 |
| | | | Þ | Þ | Þ | Þ | 01-01-01-01-01-01-01-01-01-01-01-01-01-0 |
| | | | × | × | × | × | N1=40, N2=10 |
| | | | | | | | Fold Change: 2.51 |
| | | | | | | | P-value: 0 |
| 514 | 985 | H16568 | 293.47+/-211.85 | | | 288.53+/-212.27 | 288.53+/-212.27 |
| | | | 64.62+/-58.27 | | | 46,38+/-45,45 | 38.38+/-26.86 |
| _ | | | N1=40, N2=168 | × | × | N1=40, N2=31 | N1=40, N2=10 |
| | | | Fold Change: 3.74 | | | Fold Change: 4.69 | Fold Change: 5.41 |
| | | | P-value: 0 | | | P-value: 0 | P-value: 0 |
| 515 | 786 | H27948 | 221.34+/-117.87 | | | 221.34+/-117.87 | 221.34+/-117.87 |
| | | | 86.49+/-44.22 | | | 80.01+/-37.66 | 76.82+/-36.99 |
| | | | N1=39 N2=168 | > | × | N=29 N2=31 | M=29 N2=6 |
| | | | 007 77 60 77 | • | ; | | 110 |
| | | | Fold Change: 2.46 | | | Fold Change: 2.42 | Fold Change: 2.75 |
| | | | P-value: 0 | | | P-value: 0 | P-value: .00004 |
| 516 | 990 | H42085 | | | | | 405.54+/-244.99 |
| _ | | | | | | | 174.03+/-102.69 |
| | | | × | × | × | × | N1=40, N2=10 |
| | | | | | | | Fold Change: 2.17 |
| | | | | | | | P-value: .00064 |
| 517 | 166 | H43374 | | | | | 588.83+/-221.87 |
| | | | | | | | 314.31+/-230.39 |
| | | | × | × | × | × | N1-40, N2-10 |
| | | | | | | | Fold Change: 2,1 |
| | | | | | | | P-value: .00493 |
| 518 | 992 | H54254 | 370.85+/-679.27 | | 377.04+/-687.01 | 377.04+/-687.01 | 377.04+/-687.01 |
| | | | 33.63+/-58.23 | | 31.69+/-34.95 | 21.83+/-62.85 | 12.93+/-28.05 |
| | | | N1=40, N2=168 | × | N1-40, N2-6 | N1=40, N2=31 | N1=40, N2=10 |
| _ | | | Fold Change: 5.23 | | Fold Change: 4.88 | Fold Change: 6.53 | Fold Change: 6.25 |
| | | | P-vaine: 0 | | P-value: .00045 | P-value: 0 | P-value: 0 |

| * | Seq ID | Genbank | Normal vs All | Normal vs Malignant | Normal vs Stage I | Normal vs Stage II | Normal vs Stage III |
|-----|--------|---------|-------------------|---------------------|-------------------|--------------------|---------------------|
| 519 | 993 | H58608 | 202.11+/-137.98 | | | 201.08+/-139.62 | 201.08+/-139.62 |
| | | | 82 39+1.48 64 | | | 64 47+1-33 26 | 55 32+1.25 |
| | | | N1=40 N2=168 | * | × | N1=40 N2=31 | N1=40 N2=10 |
| | | | 207 107 107 | ¢ | 4 | 10,110,111 | OI SALES |
| | | | rold Change: 2.25 | | | Fold Change: 2.1/ | Fold Change: 5.21 |
| - | | | P-value: 0 | | | F-vame; 0 | P-vame: 0 |
| 220 | 1001 | N21030 | | | | 822.97+/-445.6 | |
| | | | | | | 356.31+/-294.65 | |
| | | | × | × | × | N1=40, N2=31 | × |
| | | | | | | Fold Change: 2.5 | |
| 2 | 1064 | N21424 | | | | F-Value: .00002 | 244 38+/-141 72 |
| 1 | | | | | | | 124 11+7 57 21 |
| | | | ; | , | ; | ; | 154.117-7.131 |
| | | | × | × | × | × | N1=40, N2=10 |
| | | | | | | | Fold Change: 2.56 |
| | | | | | | | P-value: .00005 |
| 522 | 1066 | N24987 | | | | 92.04+/-95.24 | 92.04+/-95.24 |
| | | | | | | 206.91+/-82.99 | 210.74+/-113.1 |
| | | | × | × | × | N1=40, N2=31 | N1=40, N2=10 |
| | | | | | | Fold Change: 2.71 | Fold Change: 2.65 |
| | | | | | | P-value: 0 | P-value: .00009 |
| 523 | 1067 | N25096 | | | | | 308.47+/-138.2 |
| | | | | | | | 118.5+/-63.44 |
| | | | × | × | × | × | N1=40, N2=10 |
| | | | | | | | Fold Change: 2.77 |
| 474 | 1068 | L9CSCN | 26 22+4-51 01 | | | | r-value00020 |
| | | | 217.96+/-178.06 | | | | |
| | | | N1=40 N2=169 | Þ | > | > | × |
| | | | Fold Change: 5.07 | 4 | 4 | 4 | 4 |
| | | | P-value: 0 | | | | |
| 525 | 101 | N31946 | | | | | 156,39+/-61.76 |
| | | | | | | | 332,92+/-151,29 |
| | | | × | × | × | × | N1=40, N2=10 |
| | | | | | | | Fold Change: 2.11 |
| | | | | | | | P-value: .00022 |

| N3254 N42752 SG 264-4752 N1-40, N2-168 N45224 X X X X X X X X X X X X X | # | Cog m | Conbank | Normal ve All | Normal ve Malinant | Normal we Stone I | Normal ve Stage II | Normal ve Stage III |
|--|----------|--------|-----------|-------------------|------------------------|-------------------|---------------------|---------------------|
| 1074 N45722 G3.264-47.52 1074 N45724 G3.264-47.52 1075 N45224 X X X 1076 N45329 365.774-123.03 1076 N45329 16.65.48,404 N1-40, N2-68 1079 N45809 X X X 1083 N53335 X X X X X 1083 N52086 289.334-330.5 N3-40, N2-168 N1-40, N2-168 | <u>ا</u> | OT TOO | Centralin | TAOL HINT VS ASII | Ivol mar vs ivianguant | TANKING AS DIABET | Not man 18 Stage A. | 11/61 // 61 // 61 |
| 1074 M42752 G52564-4752 G5264-4752 | 979 | 1072 | N32254 | | | | | 116.51+/-51.56 |
| 1074 N42752 G5264-44/52 M1-44/52 M | | | | | | | | 249.15+/-168.21 |
| 1074 N42752 G5254-4752 1075 N45224 X X X X X X X X X X X X X X X X X X | | | | × | × | × | × | N1=40, N2=10 |
| 1075 N4522 G3.26+4-7.52 N1-40, N2-1629.22 N N1-40, N2-1629.22 N P-value: 0 1076 N45329 365.77+1.23.03 1077 N45809 X 1083 N53355 X X X X X X X X X X X X Y N1-40, N2-168 N1-40, | | | | | | | | Fold Change: 2.03 |
| 1074 N47752 63.264-47.52 1075 N45224 X X X X X X X X X X X X X X X X X X | | | | | | | | P-value: .0051 |
| 1075 N45224 X X X X X X X X X X X X X X X X X X | 527 | 1074 | N42752 | 63.26+/-47.52 | | | 63.77+/-48.02 | |
| 1075 N45224 X X X X X X X X X X X X X X X X X X | | | | 203.51+/-259.92 | | | 251.31+/-284.54 | |
| 1075 N45224 X X X X X X X X X X X X X X X X X X | | | | N1=40 N2=168 | × | × | N1=40. N2=31 | × |
| 1075 N45224 X X X X X X X X X X X X X X X X X X | | | | Fold Change: 2.23 | : | | Fold Change: 2.74 | |
| 1075 N45224 X X X X X X X X X X X 1076 N45239 365.774-123.03 X X 1076 N45399 15.605.48.04 X 1079 N45899 X 1079 N45899 X X X X X X X X X X X X X X X X X X | | | | P-value: 0 | | | P-value: .00008 | |
| 1076 N45220 36577+4.123.03 16764-4.123.04 164.024-48.04 N1-40, X2-168. N1-40, X2-168. N1-40, X2-168. N1-40, X2-168. N1-40, X2-168. N2-40, X2-168. N5-1335 X | 828 | 1075 | N45224 | | | | | 574.83+/-196.53 |
| 1076 N45329 365.774-123.03 1076 N45329 156.05-48.04 N1-40, N2-68 1079 N45899 X X X X X X X X X X X X | | | | | | | | 208.47+/-108.26 |
| 1076 N45329 3657744123.03 1640544.8404 X 1640544.8404 X 1640544.8404 X 1079 N45809 P-vither 0 X X X 1082 N51335 X N1-04, R2-108 X N1-0 | | | | × | × | × | × | N1=40, N2=10 |
| 1076 N45320 365774-123.09 1076 N45320 164.054-68.04 N1-40, N2-168 P-0d Change 2.35 P-orthor 0 X X X X X X X X X X X X X X Y 1083 N5335 X X X X X Y Y Y Y Y Y Y Y Y Y Y Y Y Y | | | | | | | | Fold Change: 2.79 |
| 1076 N45329 365.774-123.03 1676 N45329 156.054-64.04 N1-40, N2-68 X N1-40, N2-68 X N1-40, N2-68 X N + 40, N2-68 X X X X X X X N X X N X X N X X N X X N X X N X X X N X X X N X X X N X X X X X X X X X X X X X X X X X X X | | | | | | | | P-value: .00004 |
| 1079 N48899 X X X X X X X X X X X X X X X X X X | 529 | 1076 | N45320 | 365.77+/-123.03 | | | 359,46+/-117.88 | 359.46+/-117.88 |
| 1079 N4889 1079 N4889 X X X 1082 N51335 X X X 1083 N52066 289.854/310.3 N1-40, N2-168 X N1-40, N2-168 | | | | 164.05+/-84.04 | | | 163.46+/-78.87 | 150.13+/-52.84 |
| 1079 N48809 X X X X X X X X X X X X X X X X X X X | | | | N1=40, N2=168 | × | × | N1=40, N2=31 | N1=40, N2=10 |
| 1079 N48809 X X X 1082 N51335 X X X 1083 N52066 289.834/310.5 N1-40, N2-168 X Pold Change 2.07 Pold Change 2.07 Pold Change 2.07 Pold Change 2.07 Pold Change 2.07 Pold Change 2.07 Pold Change 2.07 Pold Change 2.07 Pold Change 2.07 Pold Change 2.07 Pold Change 2.07 | | | | Fold Change: 2.35 | | | Fold Change: 2.36 | Fold Change: 2.34 |
| 1079 N46809 X X X 1082 N51335 X X X 1083 N52086 289.854-310.5 N1-40, N2-108 X N1-40, N2-108 X Pold Chapege 2.07 Pold Chapege 2.07 Pold Chapege 2.07 Pold Chapege 2.07 Pold Chapege 2.07 Pold Chapege 2.07 Pold Chapege 2.07 Pold Chapege 2.07 Pold Chapege 2.07 Pold Chapege 2.07 Pold Chapege 2.07 | | | | P-value: 0 | | | P-value: 0 | P-value: .00001 |
| X X X 1082 NS1335 X X X 1083 NS2086 289.83+/-310.5 NI-40, R2-108 X Fold Changes, 207 P-value, 00114 | 330 | 1079 | N48809 | | | 130.33+/-82.96 | | 130.33+/-82.96 |
| 1062 N51335 X X X X X X X 1062 N51335 X X X X X X X X X X X X X X X X X X | | | | | | 233.51+/-111.27 | | 249.11+/-95.4 |
| 1082 NS1335 X X X X X 1083 NS2086 289.88+/210.5 NG-0, R2-08 NG-08 | | | | × | × | N1=40, N2=6 | × | N1=40, N2=10 |
| 1082 NS1335 X X X X 1083 NS2086 289.834/-310.5 NP-410.2.7 NP-40, NP-40.8 XP-410.5 XP-40.8 XP-4 | | | | | | Fold Change: 2.03 | | Fold Change: 2.14 |
| 1082 NS1335 X X X X X 1083 NS2086 289.834-210.5 N=40, N2=0.8 R=40, N2=0.8 R=40, N2=0.8 P-40, N2= | | | | | | P-value: .00829 | | P-value: .00153 |
| X X 1083 NS2086 289.534-210.5 106.774-4.10.27 N1-40, N2-68 Fold Change: 2.07 P-value: 00114 | 531 | 1082 | N51335 | | | | 108.78+/-93.56 | |
| 1083 N52086 289.83+/-310.5 1067+/-102.7 N1-40, N2-168 Fold Change, 2.07 P-value, 2011.4 P-value, 2011.4 | | | | × | × | × | N1=40, N2=31 | × |
| 1083 NS2086 289.33+/-310.5 10677+/102.27 NI=40, N2=168 Fold Change: 2.07 Ford Change: 2.07 Ford Change: 2.07 | | | | | | | Fold Change: 2 | |
| 1083 NS2086 229384-310.5 106.774-100.27 N1-40, N2-168 X Fold Change 2JJ7 P-value, 2011.4 | | | | | | | P-value: .00003 | |
| × | 532 | 1083 | N52086 | 289.83+/-310.5 | | | 289.83+/-310.5 | 289.83+/-310.5 |
| × | | | | 106.77+/-102.27 | | | 95.72+/-73.59 | 77.86+/-63.6 |
| Fold Change: 2.07 P-value: .00114 | | | | N1=40, N2=168 | × | × | N1=40, N2=31 | N1=40, N2=10 |
| P-value: .00114 | | | | Fold Change: 2.07 | | | Fold Change: 2.16 | Fold Change: 2.55 |
| | | | | P-value: .00114 | | | P-value: .00247 | P-value: .0038 |

| * | Seq ID | Genbank | Normal vs All | Normal vs Malignant | Normal vs Stage I | Normal vs Stage II | Normal vs Stage III |
|-----|--------|---------|-------------------|---------------------|-------------------|--------------------|---------------------|
| 533 | 1084 | N52352 | | | | | 268.01+/-82.24 |
| | | | | | | | 136.43+/-64.33 |
| | | | × | × | × | × | N1=40, N2=10 |
| | | | | | | | Fold Change: 2.09 |
| | | | | | | | P-value: .0074 |
| 534 | 1085 | N56877 | 109.5+/-80.79 | | | 109.5+/-80.79 | 109.5+/-80.79 |
| | | | 309.93+/-270.27 | | | 327.79+/-348.66 | 306.18+/-259.39 |
| | | | N1=40, N2=168 | × | × | N1=40, N2=31 | N1=40, N2=10 |
| _ | | | Fold Change: 2,45 | | | Fold Change: 2.15 | Fold Change: 2.68 |
| | | | P-value: 0 | | | P-value: ,00281 | P-value: .00382 |
| 232 | 1086 | N57539 | 225.43+/-101.42 | | | 225.43+/-101.42 | 225.43+/-101.42 |
| | | | 103.31+/-100.1 | | | 106.35+/-190.86 | 89.92-1/-26.98 |
| | | | N1=39, N2=168 | × | × | N1=39, N2=31 | N1=39, N2=6 |
| | | | Fold Change: 2.37 | | | Fold Change: 2.94 | Fold Change: 2.43 |
| | | | P-value: 0 | | | P-value: 0 | P-value: .00273 |
| 536 | 1087 | N59432 | | | 237.95+/-107.53 | 237,95+/-107,53 | 237.95+/-107.53 |
| | | | | | 71.26+/-38.6 | 105.02+/-65.3 | 110.93+/-111.49 |
| | | | × | × | N1=39, N2=10 | N1=39, N2=31 | N1=39, N2=6 |
| | | | | | Fold Change: 3.32 | Fold Change: 2.39 | Fold Change: 2.61 |
| | | | | | P-value: .00159 | P-value: 0 | P-value: .00543 |
| 537 | 1088 | N62126 | | | | | 280.04+/-181.07 |
| | | | | | | | 109.11+/-74.94 |
| | | | × | × | × | × | N1=40, N2=10 |
| | | | | | | | Fold Change: 2.59 |
| | | | | | | | P-value: .00734 |
| 238 | 1089 | N63237 | 217.51+/-156.97 | | 217.51+/-156.97 | 217.51+/-156.97 | 217.51+/-156.97 |
| | | | 63.12+/-54.35 | | 52.65+/-34.21 | 52,57+/-33,35 | 24.34+/-38.62 |
| | | | N1=39, N2=168 | × | N1=39, N2=10 | N1=39, N2=31 | N1=39, N2=6 |
| | | | Fold Change: 2.55 | | Fold Change: 3.16 | Fold Change: 2.9 | Fold Change: 3.83 |
| | | | P-value: 0 | | P-value: .00096 | P-value: 0 | P-value: .00001 |
| 539 | 1090 | N63913 | 463.66+/-314.65 | | | 458.01+/-316.71 | 458.01+/-316.71 |
| | | | 88.17+/-134.91 | | | 87.05+/-105.81 | 65.18+/-127.89 |
| | | | N1=40, N2=168 | × | × | N1=40, N2=31 | N1=40, N2=10 |
| | | | Fold Change: 5.84 | | | Fold Change: 5.25 | Fold Change: 7.57 |
| | | | P-value: 0 | | | P-value: 0 | P-value: .00008 |

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|-----|--------|---------|-------------------|---------------------|-------------------|--------------------|---------------------|
| * | Seq ID | Genbank | Normal vs All | Normal vs Malignant | Normal vs Stage I | Normal vs Stage II | Normal vs Stage III |
| 540 | 1001 | N64648 | | | | 262.87+/-87.71 | 262.87+/-87.71 |
| | | | | | | 129.76+/-54.11 | 114.48+/-29.26 |
| | | | × | × | × | N1=40, N2=31 | N1=40, N2=10 |
| | | | | | | Fold Change: 2.09 | Fold Change: 2.21 |
| | | | | | | P-vaine: 0 | P-vaine: 0 |
| 541 | 1092 | L989LN | | | | 210.78+/-96.34 | |
| | | | | | | 106.18+/-57.67 | |
| | | | × | × | × | NI=39, N2=31 | × |
| | | | | | | Fold Change: 1.97 | |
| | | | , | | | P-value: 0 | |
| 542 | 1094 | N79004 | | | | | 93.29+/-75.16 |
| | | | | | | | 273.05+/-182.9 |
| | | | × | × | × | × | N1=40, N2=10 |
| | | | | | | | Fold Change: 2.79 |
| | | | | | | | P-value: .00222 |
| 543 | 1095 | N80935 | 266,86+/-83,21 | | | | 266.66+/-84.29 |
| | | | 143,94+/-87.39 | | | | 130.9+/-68.88 |
| | | | N1=40, N2=168 | × | × | × | N1=40, N2=10 |
| | | | Fold Change: 2.01 | | | | Fold Change: 2.17 |
| | | | P-value: 0 | | | | P-value: .00097 |
| 24 | 1096 | N90525 | 117.6+/-226.89 | | | 118.22+/-229.82 | |
| | | | 226.51+/-203.72 | | | 255.32+/-119.15 | |
| | | | N1=40, N2=168 | × | × | N1=40, N2=31 | × |
| | | | Fold Change: 2.23 | | | Fold Change: 2.98 | |
| Ì | | | P-value: .00001 | | | P-value: 0 | |
| 35 | 1101 | R08000 | 501.45+/-685.59 | | 502,76+/-694.51 | 502.76+/-694.51 | 502.76+/-694.51 |
| | | | 78.58+/-101 | | 79.82+/-77.61 | 57.68+/-127.95 | 65.51+/-107.84 |
| | | | N1=40, N2=168 | × | N1=40, N2=6 | N1=40, N2=31 | N1=40, N2=10 |
| | | | Fold Change: 4.96 | | Fold Change: 4.45 | Fold Change: 7.21 | Fold Change: 6.92 |
| | | | P-value: 0 | | P-value: .0049 | P-value: 0 | P-value: .00004 |
| 546 | 1102 | R11248 | | | | 114.31+/-112.68 | 114.31+/-112.68 |
| | | | | | | 295.22+/-374.37 | 538.23+/-597.4 |
| | | | × | × | × | N1=40, N2=31 | N1=40, N2=10 |
| | | | | | | Fold Change: 2.09 | Fold Change: 3.42 |
| | | | | | | P-value: .00187 | P-value: .01362 |

| Normal vs Stage III | 1112.78+/-843.96 | 123.27+/-153.07 | N1=40, N2=10 | Fold Change: 10.71 | P-value: .00001 | 267.35+/-98.92 | 143.02+/-95.35 | N1=40, N2=10 | Fold Change: 2.14 | P-value: .0067 | 200.26+/-133.86 | 6.98+/-15.78 | N1=40, N2=10 | Fold Change: 6.74 | P-value: 0 | | | × | | | | | × | | 314 37-4-220 35 | 6.84+/-26.67 | N1=40, N2=10 | Fold Change: 4.76 | P-value: 0 | 241.29+/-181.34 | 17.97+/-75.13 | N1=40, N2=10 | |
|---------------------|------------------|-----------------|---------------|--------------------|-----------------|----------------|-----------------|--------------|-------------------|-----------------|-----------------|---------------|--------------|-------------------|-----------------|---------------|-----------------|---------------|-------------------|------------|----------------|-----------------|--------------|-------------------|-----------------|---------------|--------------|-------------------|----------------|-----------------|---------------|---------------|--|
| Normal vs | 1112.78+ | 123.27+ | N1=40 | Fold Char | P-value | 267.35 | 143.02+ | N1=40 | Fold Cha | P-value | 200.26+ | 486'9 | N1=40 | Fold Cha | P-va | | | - | | | | | ., | | 214 32 | 6.84+ | NI-40 | Fold Ch | P-va | 241.29 | 17.97 | N1=40 | |
| Normal vs Stage II | 1112.78+/-843.96 | 168.78+/-236.94 | N1=40, N2=31 | Fold Change: 7.78 | P-value: 0 | 267.35+/-98.92 | 156.98+/-110.51 | N1=40, N2=31 | Fold Change: 2.05 | P-value: .00002 | 200.26+/-133.86 | 16.03+/-23.1 | N1=40, N2=31 | Fold Change: 5.96 | P-value: 0 | 92.55+/-48.64 | 218.34+/-122.05 | N1=39, N2=31 | Fold Change: 2.14 | P-value: 0 | 703.42+/-425.8 | 319.29+/-199.18 | N1=40, N2=31 | Fold Change: 2.32 | 214 32±/229 35 | 37 79+/-84 48 | N1=40, N2=31 | Fold Change: 3.62 | P-value: 0 | 241.29+/-181.34 | 25.72+/-39.39 | N1=40, N2=31 | |
| Normal vs Stage I | 1112,78+/-843.96 | 257.86+/-243.05 | N1=40, N2=6 | Fold Change: 4.54 | P-value: .00794 | | | × | | | 200.26+/-133.86 | 49.74+/-62.98 | N1=40, N2=6 | Fold Change: 4.34 | P-value: .00827 | | | × | | | | | × | | 214 22±/_220 35 | 30 80+/-23 04 | N1=40, N2=6 | Fold Change: 3.38 | P-value: .0002 | | | × | |
| Normal vs Malignant | | | × | | | | | × | | | | | × | | | | | × | | | | | × | | | | × | | | | | × | |
| Normal vs All | 1107.23+/-833.81 | 264.46+/-256.11 | N1=40, N2=168 | Fold Change: 4.88 | P-value: 0 | | | × | | | | | × | | | 92.55+/-48.64 | 218.35+/-108.48 | N1=39, N2=168 | Fold Change: 2.17 | P-value: 0 | | | × | | 211 101/ 227 25 | 53.064/.01.11 | N1=40 N2=168 | Rold Change 2.91 | P-value: 00001 | 241.03+/-179.01 | 33.3+/-86.44 | N1=40, N2=168 | |
| Genbank | R20784 | | | | | R49392 | | | | | R54660 | | | | | R62346 | | | | | R67627 | | | | 102070 | MO2304 | | | | R70255 | | | |
| Seq ID | 1104 | | | | | 1100 | | | | | 1112 | | | | | 1113 | | | | | 1114 | | | | 444.0 | crit | | | | 9111 | | | |
| # | 547 | | | | | 548 | : | | | | 549 | | | | | 8 | | | | | 551 | | | | 200 | 700 | | | | 553 | | | |

| Sed ID | Genbank | Normal vs All | Normal vs Malignant | Normal vs Stage I | Normal vs Stage II | Normal vs Stage III |
|--------|---------|-------------------|---------------------|-------------------|--------------------|---------------------|
| | R73518 | 397.68+/-234.48 | | | 397.68+/-234.48 | 397.68+/-234.48 |
| | | 142.17+/-86.32 | | | 129.97+/-75.3 | 130.92+/-91.05 |
| | | N1=39, N2=168 | × | × | NI=39, N2=31 | N1=39, N2=6 |
| | | Fold Change: 2.61 | | | Fold Change: 2.92 | Fold Change: 2.93 |
| | | P-value: 0 | | | P-value: 0 | P-value: .00003 |
| | R74561 | | | | 425.23+/-350.96 | |
| | | | | | 871.35+/-705.04 | |
| | | × | × | × | N1=40, N2=31 | × |
| | | | | | Fold Change: 2.06 | |
| | | | | | P-value: .00036 | |
| 1 | R83604 | 294.9+/-858.81 | | 304.76+/-867.74 | 304.76+/-867.74 | 304.76+/-867.74 |
| | | -49.34+/-85.75 | | -70,76+/-37,34 | -62.65+/-38.9 | 42.74+/-57.17 |
| | | N1=40, N2=168 | × | N1=40, N2=6 | N1=40, N2=31 | N1-40, N2-10 |
| | | Fold Change: 3.18 | | Fold Change: 3.74 | Fold Change: 3.74 | Fold Change: 3.25 |
| | | P-value: .00005 | | P-value: .00001 | P-value: .00001 | P-value: .0002 |
| | T16144 | | | | 67.72+/-60.08 | 67.72+/-60.08 |
| | | | | | 246.32+/-239.25 | 213.1+/-108.52 |
| | | × | × | × | N1=40, N2=31 | N1=40, N2=10 |
| | | | | | Fold Change: 3.16 | Fold Change: 3.43 |
| | | | | | P-value: 0 | P-value: .00004 |
| 1 | TS7042 | | | | 286.11+/-193.74 | 286.11+/-193.74 |
| | | | | | 114.39+/-69.74 | 125.4+/-63.01 |
| | | × | × | × | N1=40, N2=31 | N1=40, N2=10 |
| | | | | | Fold Change: 2.34 | Fold Change: 2.01 |
| | | | | | P-value: 0 | P-value: ,00122 |
| 1 | T57670 | | | | 404.19+/-128.46 | 404.19+/-128.46 |
| | | | | | 212.97+/-107.98 | 186+/-108.03 |
| | | × | × | × | N1=40, N2=31 | N1=40, N2=10 |
| | | | | | Fold Change: 2.03 | Fold Change: 2.39 |
| | | | | | P-value: 0 | P-value: .00117 |
| 1 | T57773 | | | | | 214.22+/-119.35 |
| | | | | | | 76.17+/-87.27 |
| | | × | × | × | × | N1=40, N2=10 |
| | | | | | | Fold Change: 3.27 |
| | | | | | | P-value: .002 |

| Sed ID | Genbank | Normal vs All | Normal vs Malignant | Normal vs Stage I | Normal vs Stage II | Normal vs Stage III |
|--------|---------|-------------------|---------------------|-------------------|--------------------|---------------------|
| 1125 | T61106 | 164.66+/-104.9 | | | 164.66+/-104.9 | 164.66+/-104.9 |
| | | 345.33+/-209.56 | | | 339.88+/-198.04 | 341.68+/-227.02 |
| | | N1=39, N2=168 | × | × | N1=39, N2=31 | N1=39, N2=6 |
| | | Fold Change: 2.21 | | | Fold Change: 2.11 | Fold Change: 2.17 |
| | | P-value: .00001 | | | P-value: .00077 | P-value: .01513 |
| 1126 | T64447 | 216.31+/-152.39 | | 216,31+/-152,39 | 216.31+/-152.39 | 216.31+/-152.39 |
| | | 41.01+/-92.12 | | 39.2+/-138.62 | 17.71+/-47.39 | -19.08+/-58.11 |
| | | N1=39, N2=168 | × | N1=39, N2=10 | N1=39, N2=31 | N1=39, N2=6 |
| | | Fold Change: 3.99 | | Fold Change: 3.8 | Fold Change: 5.18 | Fold Change: 7.39 |
| | | P-value: 0 | | P-value: .01694 | P-value: 0 | P-value: 0 |
| 1130 | T79945 | 266,46+/-143.28 | | | | |
| | | 196.1+/-275.3 | | | | |
| | | N1=40. N2=168 | × | × | × | × |
| | | Fold Change: 2,04 | | | | |
| | | P-value: 0 | | | | |
| 1133 | T92947 | 370.07+/-299.38 | | | 359.82+/-296.09 | 359.82+/-296.09 |
| | | 173.72+/-164.88 | | | 134.2+/-88.77 | 113.03+/-41.6 |
| | | N1=40, N2=168 | × | × | N1=40, N2=31 | NI=40, N2=10 |
| | | Fold Change: 2,08 | | | Fold Change: 2.39 | Fold Change: 2.36 |
| | | P-value: .00004 | | | P-value: .00007 | P-value: .00008 |
| 1134 | T93570 | | | | 344.83+/-197.39 | 344.83+/-197.39 |
| | | | | | 156.26+/-76.45 | 137.27+/-55.02 |
| | | × | × | × | N1=40, N2=31 | N1=40, N2=10 |
| | | | | | Fold Change: 2.12 | Fold Change: 2.27 |
| | | | | | P-value: 0 | P-value: .0001 |
| 1170 | W02608 | 83.49+/-56.33 | | | 82.66+/-56.82 | 82.66+/-56.82 |
| | | 238.33+/-117.55 | | | 216.35+/-112.51 | 206.9+/-95.19 |
| | | N1=40, N2=168 | × | × | N1-40, N2-31 | N1=40, N2=10 |
| | | Fold Change: 3.06 | | | Fold Change: 2.77 | Fold Change: 2.61 |
| | | P-value: 0 | | | P-value: 0 | P-value: .00124 |
| 1171 | W02823 | 220.34+/-88.04 | | | 217.4+/-87.18 | 217.4+/-87.18 |
| | | 83,44+/-86.86 | | | 71.46+/-47.57 | 107.28+/-72.66 |
| | | N1=40, N2=168 | × | × | N1=40, N2=31 | N1=40, N2=10 |
| | | Fold Change: 2.96 | | | Fold Change: 3.16 | Fold Change: 2.32 |
| | | V | | | | 10000 |

| п | | _ | _ | _ | | _ | | _ | _ | | _ | | | ~ | | | _ | _ | _ | | Γ | | | | | | | _ | _ | _ | Γ | _ | | 4 |
|-----------------------|-----------------|-----------------|---------------|-------------------|-----------------|------------------|-----------------|--------------|-------------------|-----------------|----------------|-----------------|--------------|-------------------|-----------------|-----------------|----------------|--------------|-------------------|------------|---------------|-----------------|--------------|-------------------|-----------------|-----------------|-----------------|---------------|-------------------|------------|----------------|--------------|--------------|--------------------------------------|
| Normal vs Stage III | 299.21+/-164.12 | 49.97+/-36.95 | N1=40, N2=10 | Fold Change: 5.11 | P-value: 0 | 1139,71+/-444.58 | 291.49+/-262.25 | NI=40. NZ=10 | Fold Change: 5.29 | P-value: .00046 | 172.41+/-85.84 | 443.37+/-353.68 | N1=40, N2=10 | Fold Change: 2.22 | P-value: .00917 | | | × | | | | | × | | | 720.17+/-951.89 | 39.22+/-65.55 | N1=40, N2=10 | Fold Change: 7.71 | P-value: 0 | 100.04+/-68.17 | 213+/-119.61 | N1=40, N2=10 | Fold Change: 2.14 P-value: .00375 |
| Normal vs Stage II | 299.21+/-164.12 | 56.2+/-41.66 | N1=40, N2=31 | Fold Change: 4.81 | P-value: 0 | 1139.71+/-444.58 | 244.74+/-245.86 | N1=40. N2=31 | Fold Change: 5.59 | P-value: 0 | | | × | | | 314.68+/-111.87 | 147.05+/-53.04 | N1=39, N2=31 | Fold Change: 2.14 | P-value: 0 | 67.04+/-84.41 | 322,95+/-770,03 | N1=40, N2=31 | Fold Change: 2.05 | P-value: .00441 | 720.17+/-951.89 | 169.52+/-503.24 | N1=40, N2=31 | Fold Change: 4.62 | P-value: 0 | | | × | |
| Normal vs Stage I | 299,21+/-164.12 | 108.65+/-140.45 | N1=40, N2=6 | Fold Change: 3.51 | P-value: .03256 | | | × | | | | | × | | | | | × | | | | | × | | | | | × | | | | | × | |
| Normal vs Malignant | | | × | | | | | × | | | | | × | | | | | × | | | | | × | | | | | × | | | | | × | |
| Normal vs All | 301.61+/-162.72 | 67.74+/-52.69 | N1=40, N2=168 | Fold Change: 4.19 | P-value: 0 | 1144.81+/-440.02 | 379.83+/-343.79 | N1=40 N2=168 | Fold Change: 3.79 | P-value: 0 | | | × | | | | | × | | | | | × | | | 707.63+/-942.95 | 144.29+/-433.7 | N1=40, N2=168 | Fold Change: 5.42 | P-value: 0 | | | × | |
| Genbank | W07043 | | | | | W07304 | | | | | W22264 | | | | | W28281 | | | | | W31919 | | | | | W32480 | | | | | W55924 | | | |
| Seq ID | 1173 | | | | | 1174 | | | | | 1177 | | | | | 1181 | | | | | 1182 | | | | | 1183 | | | | | 1189 | | | |
| # | 268 | | | | | 569 | | | | | 570 | | _ | | | 571 | | | | | 572 | : | | | | 573 | | | | | 574 | | | |

| # | Seq ID | Genbank | Normal vs All | Normal vs Malignant | Normal vs Stage I | Normal vs Stage II | Normal vs Stage III |
|-----|--------|---------|-------------------|---------------------|-------------------|--------------------|---------------------|
| 575 | 1195 | W72062 | 381.86+/-163.16 | | | 380.68+/-165.12 | 380.68+/-165.12 |
| | | | 182.13+/-62.01 | | | 175.84+/-50.18 | 176.3+/-66.81 |
| | | | N1=40, N2=168 | × | X | N1=40, N2=31 | N1=40, N2=10 |
| | | | Fold Change: 2.03 | | | Fold Change: 2.04 | Fold Change: 2.07 |
| | | | · P-value: 0 | | | P-value: 0 | P-value: .00001 |
| 576 | 1196 | W72182 | | | | | 83.09+/-49 |
| | | | | | | | 231.34+/-134.05 |
| | | | × | × | × | × | N1=40, N2=10 |
| | | | | | | | Fold Change: 2.71 |
| | | | | | | | P-value: .00094 |
| 577 | 1199 | W72347 | 367.98+/-155.29 | | | 368.08+/-157.32 | 368.08+/-157.32 |
| | | | 146.12+/-193.47 | | | 94.65+/-155.82 | 261.92+/-468.76 |
| | | | N1=40, N2=168 | × | × | N1=40, N2=31 | N1=40, N2=10 |
| | | | Fold Change: 3.46 | | | Fold Change: 5.14 | Fold Change: 3.22 |
| | | | P-value: 0 | | | P-value: 0 | P-value: .03608 |
| 278 | 1200 | W72407 | 235.27+/-157.67 | | | 234.77+/-159.7 | |
| | | | 63,55+/-76.04 | | | 85.52+/-101.27 | |
| | | | N1=40, N2=168 | × | × | N1=40, N2=31 | × |
| | | | Fold Change: 3.77 | | | Fold Change: 3.02 | |
| | | | P-value: 0 | | | P-value: .00001 | |
| 579 | 1201 | W72511 | 995.7+/-434.28 | | | 988.5+/-437.53 | 988.5+/-437.53 |
| | | | 430.04+/-283.24 | | | 418.47+/-331.96 | 250.59+/-208.45 |
| _ | | | N1=40, N2=168 | × | × | N1=40, N2=31 | N1=40, N2=10 |
| | | | Fold Change: 2.53 | | | Fold Change: 2.63 | Fold Change: 4.54 |
| | | | P-value: 0 | | | P-value: 0 | P-value: .00008 |
| 280 | 1203 | W73230 | 526.33+/-307.22 | | | 524.48+/-311.01 | 524.48+/-311.01 |
| | | | 205.2+/-108.64 | | | 185.59+/-79.06 | 164.36+/-67.2 |
| | | | N1=40, N2=168 | × | × | N1=40, N2=31 | N1=40, N2=10 |
| | | | Fold Change: 2.58 | | | Fold Change: 2.72 | Fold Change: 3 |
| | | | P-value: 0 | | | P-value: 0 | P-value: 0 |
| 581 | 1204 | W73386 | 242.96+/-399.57 | 469.37+/-905.14 | 248.29+/-403.35 | 248.29+/-403.35 | 248.29+/-403.35 |
| | | | 16.99+/-76.53 | 95.29+/-88.61 | 3.09+/-55.26 | -16.75+/-32.9 | -2.29+/-42.09 |
| _ | | | N1=40, N2=168 | N1=17, N2=49 | N1=40, N2=6 | N1=40, N2=31 | N1=40, N2=10 |
| | | | Fold Change: 4.25 | Fold Change: 3.43 | Fold Change: 4.79 | Fold Change: 5.86 | Fold Change: 4.92 |
| | | | P-value: 0 | P-value: .00019 | P-value: .00079 | P-value: 0 | P-value: 0 |

| | Seq ID | Genbank | Normal vs All | Normal vs Malignant | Normal vs Stage I | Normal vs Stage II | Normal vs Stage III |
|-----|--------|---------|-------------------|---------------------|-------------------|--------------------|---------------------|
| 282 | 1205 | W73819 | | | | 993.58+/-625.18 | 993.58+/-625.18 |
| | | | | | | 399.14+/-205.47 | 277.46+/-140.58 |
| | | | × | × | × | N1=40, N2=31 | N1=40, N2=10 |
| | | | | | | Fold Change: 2.23 | Fold Change: 3.36 |
| | | | | | | P-value: 0 | P-value: .00039 |
| 583 | 1206 | W73855 | | | | 201.45+/-105.43 | |
| | | | | | | 87.02+/-86.49 | |
| | | | × | × | × | N1=40, N2=31 | × |
| | | | | | | Fold Change: 2.72 | |
| | | | | | | P-value: 0 | |
| 584 | 1207 | W73890 | 223.3+/-129.09 | | | 223.01+/-130.77 | 223.01+/-130.77 |
| | | | 73.33+/-44.79 | | | 65.62+/-33.81 | 62.68+/-63.75 |
| | | | N1=40, N2=168 | × | X | N1=40, N2=31 | N1=40, N2=10 |
| | | | Fold Change: 2.96 | | | Fold Change: 3.14 | Fold Change: 3.85 |
| | | | P-value: 0 | | | P-value: 0 | P-value: .00019 |
| 585 | 1209 | W80496 | 221.23+/-140,42 | | | 214,3+/-135,15 | 214.3+/-135.15 |
| | | | 105,98+/-75,28 | | | 95.94+/-55.75 | 53.76+/-58.28 |
| | | | N1=40, N2=168 | × | X | N1=40, N2=31 | N1=40, N2=10 |
| | | | Fold Change: 2.11 | | | Fold Change; 2.13 | Fold Change: 3.87 |
| | | | P-value: 0 | | | P-value: .00004 | P-value: .00029 |
| 286 | 1211 | W88427 | 669.14+/-243.38 | | | 661.21+/-241.27 | 661.21+/-241.27 |
| | | | 316.73+/-233.88 | | | 304.22+/-202.92 | 187.14+/-59.43 |
| | | | N1=40, N2=168 | × | × | N1-40, N2-31 | N1=40, N2=10 |
| | | | Fold Change: 2.35 | | | Fold Change: 2.35 | Fold Change: 3.45 |
| | | | P-value: 0 | | | P-value: 0 | P-value: 0 |
| 587 | 1246 | Z99386 | 615.03+/-208.26 | | | 611.71+/-209.91 | 611.71+/-209.91 |
| | | | 262.6+/-122.89 | | | 245.54+/-120.4 | 231.83+/-120.93 |
| | | | N1=40, N2=168 | × | × | N1=40, N2=31 | N1=40, N2=10 |
| | | | Fold Change: 2.48 | | | Fold Change: 2.66 | Fold Change: 3.04 |
| | | | P-value: 0 | | | P-value- 0 | D-volue: 00124 |

226 Table 5: BREAST / INFILTRATING LOBULAR CARCINOMA

| # | Seq ID | Genbank | Normal vs All | Normal vs Malignant | Normal vs SII and SIII |
|----|--------|-------------|-------------------|---------------------|--------------------------------|
| 1 | 7 | AA017070 | 218.33+/-195.52 | | • |
| | | | 77.15+/-79.13 | | |
| | | | N1=40, N2=17 | X | X |
| | | | Fold Change: 2.53 | | |
| | | | P-value: .00187 | | |
| 2 | 15 | AA031790 | 336.45+/-181.35 | | |
| | | | 156.08+/-81.33 | | |
| | | | N1=40, N2=17 | X | x |
| | | | Fold Change: 2.16 | | |
| | | | P-value: .00003 | | |
| 3 | 23 | AA044830 | 387.92+/-190.91 | • | |
| | | | 188.55+/-88.55 | | |
| | | | N1=40, N2=17 | X | x |
| | | | Fold Change: 2.14 | | |
| | | | P-value: .00023 | | |
| 4 | 24 | AA045145 | 262.21+/-180.28 | | |
| | | | 76.07+/-123.14 | | NP. |
| | | | N1=40, N2=17 | X | x |
| | | | Fold Change: 3.26 | | |
| | | | P-value: .00038 | | |
| 5 | 25 | AA046457 | 254.96+/-154.86 | | |
| | | | 128.89+/-118.57 | | |
| | | | N1=40, N2=17 | X | X |
| | | | Fold Change: 2.3 | | |
| | | | P-value: .00176 | | |
| 6 | 31 | AA059396 | 383.25+/-127.97 | | 383.25+/-127.97 |
| | | | 170.7+/-70.05 | | 120.28+/-48.53 |
| | | | N1=40, N2=17 | X | N1=40, N2=17 |
| | | | Fold Change: 2.32 | | Fold Change: 3.22 |
| | | | P-value: 0 | | P-value: .01218 |
| 7 | 33 | AA059458 | 74.76+/-90.6 | | 74.76+/-90.6 |
| | | | 314.12+/-111.83 | x | 344.29+/-46.75 N1=40, N2=17 |
| | | | N1=40, N2=17 | X | Fold Change: 6.82 |
| | | | Fold Change: 5.79 | | P-value: 0 |
| | | | P-value: 0 | | r-value: 0 |
| 8 | 41 | AA126704 | 312.64+/-137.34 | | |
| | | | 130.96+/-82.96 | v | x |
| | | | N1=40, N2=17 | X | ^ |
| | | | Fold Change: 2.5 | | |
| | | 1.1.1.00011 | P-value: .00009 | | |
| 9 | 42 | AA127718 | 240.21+/-361.64 | | |
| | | | 75.73+/-121.03 | x | x |
| | | | N1=40, N2=17 | X | ^ |
| | | | Fold Change: 3.09 | | |
| | | | P-value: .00005 | | |
| 10 | 43 | AA127727 | 212.97+/-123.48 | | |
| | | | 100.07+/-53.82 | 32 | x |
| | | | N1=40, N2=17 | X | |
| | | | Fold Change: 2.1 | | |
| | | | P-value: .00014 | | |
| 11 | 51 | AA133248 | 400.91+/-134.73 | | |
| | | | 201.52+/-119.8 | •• | |
| | | | N1=40, N2=17 | X | X |
| | | | Fold Change: 2.24 | | |
| | | | P-value: .00009 | | |

| # | Seq ID | Genbank | Normal vs All | Normal vs Malignant | Normal vs SII and SIII |
|----|--------|-----------|-------------------|---------------------|------------------------|
| 12 | 57 | AA142913 | 302.34+/-222.83 | | 302.34+/-222.83 |
| | | | 104.53+/-62.4 | | 68.29+/-31.77 |
| | | | N1=40, N2=17 | X | N1=40, N2=17 |
| | | | Fold Change: 2.84 | | Fold Change: 4.03 |
| | | | P-value: 0 | | P-value: .00871 |
| 13 | 62 | AA147751 | 478.2+/-207.42 | | |
| 13 | 02 | AMITTION. | 245,52+/-144,78 | | |
| | | | N1=40, N2=17 | x | X |
| | | | Fold Change: 2.03 | | |
| | | | P-value: .00015 | | |
| 14 | 63 | AA147884 | 46.86+/-55.16 | | |
| 14 | 0.3 | AA14/004 | 212.3+/-151.24 | | |
| | | | | x | x |
| | | | N1=40, N2=17 | | Α. |
| | | | Fold Change: 3.93 | | |
| | | | P-value: .00001 | .,, | |
| 15 | 64 | AA149312 | 374+/-139.43 | | |
| | | | 179.7+/-77.1 | | 37 |
| | | | N1=40, N2=17 | X | X |
| | | | Fold Change: 2.18 | | |
| | | | P-value: .00003 | | |
| 16 | 65 | AA150501 | 215.8+/-104 | | |
| | | | 97.75+/-48.53 | | |
| | | | N1=40, N2=17 | X | X |
| | | | Fold Change: 2.27 | | |
| | | | P-value: .00006 | | |
| 17 | 71 | AA158731 | 287.72+/-241.22 | | |
| | | | 94.76+/-99 | | |
| | | | N1=40, N2=17 | x | x |
| | | | Fold Change: 3,29 | | |
| | | | P-value: .00036 | | |
| 18 | 72 | AA160156 | 630.23+/-274.77 | | |
| | | | 297.85+/-166.73 | | |
| | | | N1=40, N2=17 | X | x |
| | | | Fold Change: 2.39 | | |
| | | | P-value: .00076 | | |
| 19 | 75 | AA173572 | 368.73+/-173.58 | | 368.73+/-173.58 |
| | | 122210012 | 140.6+/-66.1 | | 101.84+/-30.25 |
| | | | N1=40, N2=17 | x | N1=40, N2=17 |
| | | | Fold Change: 2.52 | | Fold Change: 3.17 |
| | | | P-value: .00001 | | P-value: .00053 |
| 20 | 84 | AA203663 | 288.39+/-92.75 | | |
| 20 | 04 | 242203003 | 151.54+/-90.12 | | |
| | | | N1=40, N2=17 | x | x |
| | | | Fold Change: 2.19 | * | |
| | | | P-value: .00062 | | |
| 21 | 88 | AA227778 | 254.32+/-164.5 | | |
| 41 | 80 | AA44///8 | 129,32+/-121.52 | | |
| | | | | x | x |
| | | | N1=40, N2=17 | | |
| | | | Fold Change: 2.21 | | |
| | | | P-value: .00551 | | |
| 22 | 99 | AA369887 | 326.24+/-259.48 | | |
| | | | 1569.71+/-1564.61 | | |
| | | | N1=40, N2=17 | x | X |
| | | | Fold Change: 3.13 | | |
| | | | P-value: .00723 | | |

| # | Seq ID | Genbank | Normal vs All | Normal vs Malignant | Normal vs SII and SIII |
|------|--------|----------|-------------------|---------------------|------------------------|
| 23 | 117 | AA430314 | 259.57+/-186.05 | | 259.57+/-186.05 |
| | | | 94.12+/-84.62 | | 51.43+/-32.49 |
| | | | N1=40, N2=17 | х . | N1=40, N2=17 |
| | | | Fold Change: 2.81 | | Fold Change: 4.33 |
| | | | P-value: .00057 | | P-value: .0109 |
| 24 | 120 | AA447015 | 226.67+/-173.74 | | 226.67+/-173.74 |
| ٦. | | | 86,47+/-87,06 | | 49.75+/-49 |
| 1 | | | N1=40, N2=17 | x | N1=40, N2=17 |
| | | | Fold Change: 2.44 | ·- | Fold Change: 3.69 |
| | | | P-value: .00239 | | P-value: .04932 |
| 25 | 121 | AA448195 | 82,22+/-92.11 | | |
| | | | 252.38+/-226.28 | | |
| 1 | | | N1=40, N2=17 | X | x |
| 1 | | | Fold Change: 2.56 | | |
| | | | P-value: .00561 | | |
| 26 | 122 | AA450090 | 285.47+/-226.15 | | 285.47+/-226.15 |
| | | | 121.51+/-105.64 | | 80.74+/-67.6 |
| | | | N1=40, N2=17 | x | N1=40, N2=17 |
| | | | Fold Change: 2.56 | | Fold Change: 3.67 |
| | | | P-value: .00017 | | P-value: .04277 |
| 27 | 124 | AA452295 | 220.36+/-116.43 | | 220.36+/-116.43 |
| -' | | | 43.55+/-34.23 | | 27.93+/-7.95 |
| | | | N1=40, N2=17 | X | N1=40, N2=17 |
| | | | Fold Change: 4.8 | | Fold Change: 6.64 |
| | | | P-value: 0 | | P-value: 0 |
| 28 | 129 | AA479033 | 105,96+/-264.08 | | |
| - " | | | 699.96+/-1244.37 | | |
| | | | N1=40, N2=17 | x | X |
| | | | Fold Change: 3.25 | | |
| | | | P-value: .01862 | | |
| 29 | 131 | AA480075 | 331.5+/-159.34 | | |
| T . | | | 170.51+/-174.22 | | |
| | | | N1=40, N2=17 | x | x |
| | | | Fold Change: 2.36 | | |
| | | | P-value: .00065 | | |
| 30 | 134 | AA486731 | 417.18+/-216.76 | | |
| - ' | | | 258.38+/-279.38 | | |
| | | | N1=40, N2=17 | x | x |
| | | | Fold Change: 2.26 | | |
| | | | P-value: .0077 | | |
| 31 | 135 | AA488889 | 298.86+/-194.94 | | |
| | | | 114.61+/-41.42 | | |
| | | | N1=40, N2=17 | X | x |
| | | | Fold Change: 2.16 | | |
| | | | P-value: .00001 | | |
| 32 | 138 | AA502943 | 439.24+/-110.96 | | |
| | | | 200.97+/-110.89 | | |
| | | | N1=40, N2=17 | X | X |
| | | | Fold Change: 2.41 | | |
| | | | P-value: 0 | | |
| 33 | 140 | AA508196 | 475.57+/-315.6 | | |
| | 4 | | 208.59+/-128.6 | | |
| | | | N1=40, N2=17 | x | x |
| | | | Fold Change: 2,29 | • | |
| | | | P-value: .0014 | | |
| L.—— | | | | | |

| # | Seq ID | Genbank | Normal vs All | Normal vs Malignant | Normal vs SII and SIII |
|------|--------|-----------|-------------------|---------------------|------------------------------|
| 34 | 142 | AA516420 | 208.7+/-209.98 | | |
| | | | 762.28+/-919.5 | | |
| | | | N1=40, N2=17 | x | X |
| | | | Fold Change: 2.83 | | |
| | | | P-value: .00199 | | |
| 35 | 151 | AA526961 | 417.14+/-237.24 | | |
| | | | 139.33+/-66.58 | | |
| | | | N1=40, N2=17 | x | X |
| | | | Fold Change: 2.89 | | |
| | | | P-value: 0 | | |
| 36 | 156 | AA534456 | 1130.9+/-759.82 | | |
| | | | 504.53+/-276.17 | | |
| | | | N1=40, N2=17 | x | x |
| | | | Fold Change: 2.23 | | |
| | | | P-value: .00282 | | |
| 37 | 160 | AA535218 | 322.09+/-137.43 | | |
| 31 | 100 | 24.555210 | 130.51+/-83.58 | | |
| | | | N1=40, N2=17 | x | x |
| | | | Fold Change: 2.69 | * | |
| | | | P-value: .00001 | | |
| 38 | 171 | AA584310 | 402.55+/-323.55 | | |
| 30 | 1/1 | AA304310 | 1185.08+/-725.81 | | |
| | | | N1=40, N2=17 | x | x |
| | | | Fold Change: 3.27 | Α. | |
| | | | | | |
| | | | P-value: .00003 | | 593.26+/-1291.79 |
| 39 | 172 | AA584403 | 593.26+/-1291.79 | | 46.94+/-41.5 |
| | | | 73.69+/-113.44 | x | 46.94+7-41.5 N1=40, N2=17 |
| | | | N1=40, N2=17 | | Fold Change: 4.08 |
| | | | Fold Change: 3.63 | | P-value: .01967 |
| | | | P-value: .0001 | | P-value; .01967 |
| 40 | 175 | AA601511 | 2941.11+/-4823.41 | | |
| | | | 8196.8+/-10494.86 | ** | 37 |
| | | | N1=40, N2=17 | X | X |
| | | | Fold Change: 3.59 | | |
| | | | P-value; .04627 | | |
| 41 | 178 | AA609310 | 285.39+/-160.8 | | • |
| | | | 103.37+/-63.8 | | |
| | | | N1=40, N2=17 | X | X |
| | | | Fold Change: 2.73 | | |
| | | | P-value: .00003 | | |
| 42 | 180 | AA610522 | 803+/-768.74 | | 803+/-768.74 |
| | | | 2236.91+/-2047.57 | | 1948.9+/-1536.5 |
| | | | N1=40, N2=17 | x | N1=40, N2=17 |
| | | | Fold Change: 3.15 | | Fold Change: 3.65 |
| | | | P-value: .00504 | | P-value: .04632 |
| 43 | 184 | AA621478 | 398.69+/-325.12 | | |
| | | | 105.85+/-99.55 | | |
| | | | N1=40, N2=17 | X | X |
| | | | Fold Change: 3.76 | | |
| | | | P-value: .00002 | | |
| 44 | 189 | AA628467 | 1145.06+/-502.33 | | 1145.06+/-502.33 |
| -7-4 | 107 | AMAGE 101 | 483,55+/-276,22 | | 263.82+/-233.17 |
| | | | N1=40, N2=17 | x | N1=40, N2=17 |
| | | | Fold Change: 2.59 | Λ. | Fold Change: 5.48 |
| | | | P-value: .00016 | | P-value: .04561 |

| # | Seq ID | Genbank | Normal vs All | Normal vs Malignant | Normal vs SII and SIII |
|-----|--------|-------------|-------------------|---------------------|------------------------|
| 45 | 191 | AA631047 | 615.9+/-364.24 | | |
| | | | 335.52+/-248.64 | | |
| | | | N1=40, N2=17 | x | x |
| | | | Fold Change: 2.12 | | |
| | | | P-value: .00214 | | |
| 46 | 194 | AA634799 | 739,38+/-608,62 | | |
| -10 | | 12.200 1155 | 265.99+/-273.02 | | |
| | | | N1=40, N2=17 | x | x |
| | | | Fold Change: 3.37 | | |
| | | | P-value; .00153 | | |
| 47 | 198 | AA669106 | 84.29+/-131.22 | | |
| ••• | 2,0 | 122007200 | 224.41+/-230.31 | | |
| | | | N1=40, N2=17 | x | x |
| | | | Fold Change: 3.18 | | |
| | | | P-value: .00001 | | |
| 48 | 200 | AA700621 | 467.51+/-455.09 | | 467.51+/-455.09 |
| 10 | 200 | 72/2/00022 | 127.5+/-198.7 | | 65.41+/-73.63 |
| | | | N1=40, N2=17 | x | N1=40, N2=17 |
| | | | Fold Change: 3.36 | | Fold Change: 4.6 |
| | | | P-value: .00047 | | P-value: .03306 |
| 49 | 214 | AA742697 | 1026.03+/-1071.41 | | 1026.03+/-1071.41 |
| 77 | 214 | 242172021 | 497.89+/-1362.07 | | 72.76+/-23.65 |
| | | | N1=40, N2=17 | x | N1=40, N2=17 |
| | | | Fold Change: 3.28 | 24 | Fold Change: 7.24 |
| | | | P-value: .00238 | | P-value: 0 |
| 50 | 253 | AA921809 | 459.15+/-1266.29 | | |
| 30 | 233 | 222/2100/ | 1144,77+/-1121.05 | | |
| | | | N1=40, N2=17 | X | x |
| | | | Fold Change: 2.76 | | |
| | | | P-value: .00483 | | |
| 51 | 254 | AA921830 | 92.93+/-115.1 | | 92.93+/-115.1 |
| 21 | 254 | PARTON | 214.98+/-154.53 | | 328.17+/-235.36 |
| | | | N1=40, N2=17 | x | N1=40, N2=17 |
| | | | Fold Change: 2.53 | | Fold Change: 4.07 |
| | | | P-value: .00048 | | P-value: .03148 |
| 52 | 255 | AA921922 | 312.44+/-292.63 | | 312,44+/-292.63 |
| -54 | 200 | | 101.23+/-57.27 | | 79.08+/-33.3 |
| | | | N1=40, N2=17 | x | N1=40, N2=17 |
| | | | Fold Change: 2.73 | •• | Fold Change: 3.21 |
| | | | P-value: .00001 | | P-value: .00566 |
| 53 | 260 | AA936632 | | | 125.03+/-127.3 |
| 55 | 200 | -272/3003# | | | 341.96+/-182.6 |
| | | | x | x | N1=40, N2=17 |
| | | | A | 24 | Fold Change: 3.13 |
| | | | | | P-value: .02208 |
| 54 | 266 | AA976064 | 363.9+/-153.14 | | |
| | 200 | ALAL / GOOM | 150.7+/-67.67 | | |
| | | | N1=40, N2=17 | х | X |
| | | | Fold Change: 2.48 | | |
| | | | P-value: 0 | | |
| 55 | 281 | AC004770 | r-value, v | | 222,34+/-159,84 |
| 33 | 201 | AC004//0 | | | 51.7+/-14.58 |
| | | | x | x | N1=40, N2=17 |
| | | | | | Fold Change: 3.51 |
| | | | | | P-value: .00008 |
| | | | | | T -Autro: '00000 |

| # | Seq ID | Genbank | Normal vs All | Normal vs Malignant | Normal vs SII and SIII |
|----|--------|------------|-------------------|---------------------|------------------------|
| 56 | 297 | AF052142 | 307.17+/-169.55 | | |
| - | | | 101.76+/-54.87 | | |
| | | | N1=40, N2=17 | X | X |
| | | | Fold Change: 2.92 | | |
| | | | P-value: 0 | | |
| 57 | 317 | AI018523 | 422.08+/-187.64 | | |
| 31 | 317 | A1010343 | 137.17+/-133.59 | | |
| | | | N1=40, N2=17 | X | x |
| | | | Fold Change: 3.55 | | |
| | | | P-value: .00002 | | |
| | | AI031771 | 85.9+/-105.07 | | |
| 58 | 321 | A1031//1 | 273.11+/-256.97 | | |
| | | | N1=40, N2=17 | x | X |
| | | | | Α. | 24 |
| | | | Fold Change: 2.82 | | |
| | | | P-value: .00563 | | |
| 59 | 324 | A1039005 | 203.54+/-131.69 | | |
| l | | | 79.78+/-68.07 | ** | x |
| | | | N1=40, N2=17 | X | |
| | | | Fold Change: 2.7 | | |
| | | | P-value: .00048 | | 1005 041/14160 50 |
| 60 | 325 | AI039722 | | | 1007.24+/-1162.59 |
| | | | | | 71.46+/-83.95 |
| | | | X | X | N1=40, N2=17 |
| | | | | | Fold Change: 11.94 |
| | | | | | P-value: .00965 |
| 61 | 331 | AI057450 | 381.32+/-1572.07 | | 381.32+/-1572.07 |
| | | | -3.82+/-29.02 | | -11.17+/-8.38 |
| | | | N1=40, N2=17 | x | N1=40, N2=17 |
| | | | Fold Change: 3.3 | | Fold Change: 3.63 |
| | | | P-value: .00001 | | P-value: 0 |
| 62 | 333 | AI073394 | 124.23+/-101.36 | | |
| | | | 255.64+/-158.11 | | |
| | | | N1=40, N2=17 | x | X |
| | | | Fold Change: 2.2 | | |
| | | | P-value: .00025 | | |
| 63 | 335 | AI073992 | 110.23+/-145.3 | | |
| | | | 533.62+/-785.24 | | |
| | | | N1=40, N2=17 | . x | x |
| | | | Fold Change: 3.22 | | |
| | | | P-value: .00574 | | |
| 64 | 338 | AI079545 | 248,94+/-138.38 | | |
| " | 555 | | 465.02+/-171.05 | | |
| 1 | | | N1=40, N2=17 | x . | x |
| 1 | | | Fold Change: 2.01 | | |
| | | | P-value: .00007 | | |
| 65 | 341 | AI083598 | 339.56+/-289.33 | | 339.56+/-289.33 |
| 03 | 341 | A1003330 | 75.11+/-72.52 | | 38.38+/-30.41 |
| 1 | | | N1=40, N2=17 | x | N1=40, N2=17 |
| Į. | | | Fold Change: 3.79 | A | Fold Change: 5.72 |
| | | | P-value: .00003 | | P-value: .00274 |
| - | 0.40 | 1700.001.4 | 301.2+/-152.86 | | A-VIIIIO. 10027-T |
| 66 | 342 | AI086614 | | | |
| | | | 128.33+/-84.7 | x | x |
| | | | N1=40, N2=17 | X | |
| | | | Fold Change: 2.51 | | |
| 1 | | | P-value: .00041 | | |

| # | Seq ID | Genbank | Normal vs All | Normal vs Malignant | Normal vs SII and SIII |
|-----|--------|------------|-------------------|---------------------|------------------------|
| 67 | 343 | AI087975 | 68.87+/-58.02 | | |
| | | | 211.46+/-250.57 | | |
| | | | N1=40, N2=17 | X | x |
| | | | Fold Change: 2.28 | | |
| | | | P-value: .00976 | | |
| 68 | 344 | AI088609 | 709.25+/-600.21 | | |
| | | | 265.96+/-356.75 | | |
| | | | N1=40, N2=17 | X | x |
| | | | Fold Change: 3.21 | | |
| | | | P-value: .00094 | | |
| 69 | 345 | AI091154 | 351.29+/-406.17 | | 351.29+/-406.17 |
| | | | 74.97+/-110.43 | | 12.49+/-5.56 |
| | | | N1=40, N2=17 | x | N1=40, N2=17 |
| | | | Fold Change: 4.1 | | Fold Change: 8.99 |
| | | | P-value: .00011 | | P-value: 0 |
| 70 | 351 | AI123555 | 300+/-164.6 | | 300+/-164.6 |
| | | | 65.25+/-46.06 | | 48.57+/-47.56 |
| | | | N1=40, N2=17 | X | N1=40, N2=17 |
| | | | Fold Change: 4.55 | | Fold Change: 6 |
| | | | P-value: 0 | | P-value: .01993 |
| 71 | 359 | AI128820 | 224.42+/-90.96 | | |
| | 50) | ,11120020 | 108.28+/-86.45 | | |
| | | | N1=40, N2=17 | x | x |
| | | | Fold Change: 2.34 | | |
| | | | P-value: .00033 | | |
| 72 | 361 | AI129626 | 278.92+/-134.16 | | |
| 12 | 301 | 73377020 | 134.17+/-77.75 | | |
| | | | N1=40, N2=17 | x | x |
| | | | Fold Change: 2.13 | | |
| | | | P-value: .00023 | | |
| 73 | 362 | AI131078 | 299.48+/-223.81 | | 299.48+/-223.81 |
| ,,, | 502 | PALIDIO 70 | 111.16+/-71.9 | | 67.7+/-89.93 |
| | | | N1=40, N2=17 | X | N1=40, N2=17 |
| | | | Fold Change: 2.6 | | Fold Change: 5.06 |
| | | | P-value: .0002 | | P-value: .04594 |
| 74 | 370 | AI148006 | 241.17+/-193.5 | | |
| ′- | 370 | 14440000 | 77.61+/-92.82 | | |
| | | | N1=40, N2=17 | x | x |
| | | | Fold Change: 2.93 | | |
| | | | P-value: .00043 | | |
| 75 | 372 | AI149637 | 212.6+/-241.64 | | 212.6+/-241.64 |
| ,,, | 312 | 2311-12027 | 39.92+/-27.3 | | 39.29+/-41.66 |
| | | | N1=40, N2=17 | х | N1=40, N2=17 |
| | | | Fold Change: 3.37 | | Fold Change: 3.31 |
| | | | P-value: 0 | | P-value: .04204 |
| 76 | 380 | AI189011 | 284.7+/-101.6 | | |
| /0 | 300 | A1102011 | 126.14+/-81.81 | | |
| | | | N1=40, N2=17 | х | x |
| | | | Fold Change: 2.75 | X | |
| | | | P-value: .00017 | | |
| | | 17000074 | | | |
| 77 | 384 | AI200954 | 524.84+/-319.36 | | |
| | | | 253.81+/-173.45 | x | x |
| | | | N1=40, N2=17 | X | Α. |
| | | | Fold Change: 2.17 | | |
| | | | P-value: .00291 | | |

| 78 | 386 | AI201965 | | | 234.24+/-149.37 |
|----|-----|----------|-------------------|---|--------------------|
| | | | | | |
| | | | | | 59.16+/-44.89 |
| | | | X | X | N1=40, N2=17 |
| | | | | | Fold Change: 3.61 |
| | | | | | P-value: .03602 |
| 79 | 394 | AI222594 | 431.73+/-162.38 | | |
| | | | 196.71+/-138.58 | | |
| | | | N1=40, N2=17 | X | X |
| | | | Fold Change: 2.48 | | |
| | | | P-value: .00005 | | |
| 80 | 395 | AI223817 | 221.5+/-204.3 | | |
| | | | 686.72+/-465.96 | x | х |
| | | | N1=40, N2=17 | х | X |
| | | | Fold Change: 3.28 | | |
| | | | P-value: .00041 | | |
| 81 | 399 | AI247837 | 250.33+/-314.52 | | 250.33+/-314.52 |
| | | | 53.27+/-43.26 | | 28.03+/-28.56 |
| | | | N1=40, N2=17 | x | N1=40, N2=17 |
| | | | Fold Change: 2.95 | | Fold Change: 4.49 |
| | | | P-value: .00014 | | P-value: .00427 |
| 82 | 408 | AI277612 | 1022.91+/-907.07 | | 387.19+/-203.85 |
| | | | 101.24+/-106.96 | | 584.56+/-51.28 |
| | | | N1=40, N2=17 | x | N1=40, N2=17 |
| | | | Fold Change: 8.06 | | Fold Change: 2.01 |
| | | | P-value: 0 | | P-value: .00012 |
| 83 | 417 | AI300876 | 601.83+/-985.51 | | 601.83+/-985.51 |
| | | | 26,36+/-32.43 | | 28.36+/-46.5 |
| | | | N1=40, N2=17 | x | N1=40, N2=17 |
| | | | Fold Change: 7.1 | | Fold Change: 6.7 |
| | | | P-value: 0 | | P-value: .00688 |
| 84 | 418 | AI301060 | 1095.7+/-461.79 | | |
| | | | 3285.81+/-2230.69 | | |
| | | | N1=40, N2=17 | x | x |
| | | | Fold Change: 2.58 | | |
| | | | P-value: .00018 | | |
| 85 | 422 | AI333767 | 201.68+/-104.32 | | |
| | | | 94.33+/-75 | | |
| | | | N1=40, N2=17 | x | x |
| | | | Fold Change: 2.32 | | |
| | | | P-value: .00023 | | |
| 86 | 423 | AI333987 | | | 208.53+/-320.79 |
| | | | | | -12.06+/-45.78 |
| | | | X | X | N1=40, N2=17 |
| | | | | | Fold Change: 4.29 |
| | | | | | P-value: .00037 |
| 87 | 427 | AI341602 | 137.44+/-280.1 | | 137.44+/-280.1 |
| | | | 473,63+/-503.04 | | 1084.1+/-558.85 |
| | | | N1=40, N2=17 | x | N1=40, N2=17 |
| | | | Fold Change: 3.72 | | Fold Change: 14.07 |
| | | | P-value: .00123 | | P-value: .00013 |
| 88 | 430 | AI344312 | 85.72+/-58.03 | | A |
| 00 | 750 | MISTISIA | 241.24+/-132.01 | | |
| | | | N1=40, N2=17 | X | x |
| | | | | | |
| | | | Fold Change: 2.77 | A | 22 |

| # | Seq ID | Genbank | Normal vs All | Normal vs Malignant | Normal vs SII and SIII |
|----|--------|-----------|-------------------|---------------------|--------------------------------------|
| 89 | 431 | AI346341 | 635.18+/-426.52 | | |
| | | | 192.7+/-146.21 | | |
| | | | N1=40, N2=17 | X | X |
| | | | Fold Change: 2.74 | | |
| | | | P-value: .00095 | | |
| 90 | 442 | AI369840 | 239.87+/-167.43 | | |
| | | | 91.16+/-73.21 | | |
| | | | N1=40, N2=17 | x | X |
| | | | Fold Change: 2.54 | | |
| | | | P-value: .00091 | | |
| 91 | 447 | AI378584 | 815,22+/-371.96 | | 815,22+/-371,96 |
| 71 | *** | 741570507 | 289.2+/-132.28 | | 225.35+/-105.83 |
| | | | N1=40, N2=17 | x | N1=40, N2=17 |
| | | | Fold Change: 2.65 | | Fold Change: 3.53 |
| | | | P-value: 0 | | P-value: .02945 |
| 92 | 448 | AI379723 | 380.22+/-173.64 | | |
| 92 | 440 | A13/9/43 | 171.75+/-85.82 | | |
| | | | N1=40, N2=17 | x | x |
| | | | Fold Change: 2.11 | Α. | A |
| | | | P-value: .00049 | | |
| | 450 | AI394013 | r-value. 100049 | | 81.65+/-57.28 |
| 93 | 459 | A1394013 | | | 206.8+/-28.72 |
| | | | x | x | N1=40, N2=17 |
| | | | | ^ | Fold Change: 3.01 |
| | | | | | P-value: 0 |
| | | | *** *** * *** | | 933.35+/-487.41 |
| 94 | 462 | AI417267 | 933.35+/-487.41 | | 232.02+/-44.3 |
| | | | 367.83+/-178.5 | x | |
| | | | N1=40, N2=17 | X | N1=40, N2=17 Fold Change: 3.44 |
| | | | Fold Change: 2.35 | | P-value: 0 |
| | | | P-value: 0 | | 445.97+/-259.12 |
| 95 | 467 | AI419030 | 445.97+/-259.12 | | 100.89+/-50.85 |
| | | | 141.54+/-110.04 | 37 | |
| | | | N1=40, N2=17 | X | N1=40, N2=17 |
| | | | Fold Change: 3.4 | | Fold Change: 3.94 P-value: .00968 |
| | | | P-value: .00002 | | P-value: .00968 |
| 96 | 468 | AI421837 | 293.96+/-147.73 | | |
| | | | 122.58+/-60.8 | | ** |
| | | | N1=40, N2=17 | X | X |
| | | | Fold Change: 2.25 | | |
| | | | P-value: .00003 | | |
| 97 | 477 | AI458003 | 280.16+/-202.76 | | 280.16+/-202.76 |
| | | | 58.35+/-64.44 | | 29.02+/-54.63 |
| | | | N1=40, N2=17 | x | N1=40, N2=17 |
| | | | Fold Change: 4.09 | | Fold Change: 6.1 |
| | | | P-value: 0 | | P-value: .01261 |
| 98 | 484 | AI479262 | 56.35+/-67.19 | | |
| | | | 253.01+/-258.86 | | |
| | | | N1=40, N2=17 | x | x |
| | | | Fold Change: 3.34 | | |
| | | | P-value: .00113 | | |
| 99 | 489 | AI492051 | 382,34+/-177,78 | | 382.34+/-177.78 |
| ,, | 407 | 244.52051 | 99.97+/-58.1 | | 84.79+/-58.36 |
| | | | N1=40, N2=17 | x | N1=40, N2=17 |
| | | | Fold Change: 3.83 | A | Fold Change: 4.59 |
| | | | P-value: 0 | | P-value: .01274 |

| # | Seq ID | Genbank | Normal vs All | Normal vs Malignant | Normal vs SII and SIII |
|------|--------|------------|-------------------|---------------------|------------------------|
| 100 | 493 | AI492879 | 219.42+/-658.12 | | |
| | | | 360.39+/-664.73 | | |
| | | | N1=40, N2=17 | x | x |
| | | | Fold Change: 3.18 | | |
| | | | P-value: .00218 | | _ |
| 101 | 500 | A1524085 | 388.89+/-529.52 | | |
| 101 | 500 | 11121100 | 77.76+/-117.23 | | |
| | | | N1=40, N2=17 | . x | X |
| | | | Fold Change: 3.83 | | |
| | | | P-value: .00013 | | |
| 102 | 501 | AI525044 | 316.89+/-143.08 | | |
| 102 | 301 | ALDESOTT , | 163.75+/-85.16 | | |
| | | | N1=40, N2=17 | x | x |
| | | | Fold Change: 2.13 | | |
| | | | P-value: .00114 | | |
| 103 | 505 | AI537407 | 278.8+/-204.74 | | |
| 103 | 505 | AJ55/40/ | 783.29+/-533.91 | | |
| | | | | x | x |
| | | | N1=40, N2=17 | | A |
| | | | Fold Change: 2.81 | | |
| | | | P-value: .00083 | | |
| 104 | 506 | AI539386 | 1924.9+/-2430.34 | | |
| | | | 6121.55+/-7013.05 | N/ | x |
| | | | N1=40, N2=17 | X | |
| | | | Fold Change: 3.2 | | |
| | | | P-value: .00044 | | |
| 105 | 511 | AI554514 | 90.74+/-52.8 | | |
| | | | 201.02+/-166.43 | | 37 |
| | | | N1=40, N2=17 | X | х |
| | | | Fold Change: 2.08 | | |
| | | | P-value: .00026 | | |
| 106 | 512 | AI557210 | 129.15+/-140.98 | | 129.15+/-140.98 |
| | | | 491.52+/-264.84 | | 573+/-162.6 |
| | | | N1=40, N2=17 | X | N1=40, N2=17 |
| | | | Fold Change: 5.08 | | Fold Change: 6.68 |
| | | | P-value: 0 | | P-value: .00001 |
| 107 | 517 | AI566038 | 257.62+/-109.32 | | |
| | | | 124.43+/-63.25 | | |
| | | | N1=40, N2=17 | x | x |
| | | | Fold Change: 2.16 | | |
| | | | P-value: .00015 | | |
| 108 | 520 | AI571525 | 265.11+/-78.71 | | |
| | | | 141.93+/-62.73 | | |
| | | | N1=40, N2=17 | x | x |
| | | | Fold Change: 2.04 | | |
| | | | P-value: .00015 | | |
| 109 | 536 | AI624853 | 373.05+/-166.36 | | |
| 1.37 | 200 | | 180.19+/-106.47 | | |
| | | | N1=40, N2=17 | x | x |
| | | | Fold Change: 2.21 | | |
| | | | P-value: .00004 | | |
| 110 | 540 | AI634852 | 278.07+/-162.92 | | |
| 110 | 540 | A1034834 | 122,35+/-122.97 | | |
| | | | N1=40, N2=17 | x | x |
| | | | Fold Change: 2.6 | Α. | Α. |
| | | | | | |
| | | | P-value: .00095 | | |

| # Seq ID Genbank Normal vs All Normal vs All Normal vs All and St 20,244+376.87 111 542 AI638295 | SIII |
|--|------|
| X | |
| Teld Change: 3.16 P-value: 0 | |
| 112 545 | |
| 112 545 | 6 |
| 209.61+.974.9 N=40, N2=17 X X N=40, N2=17 X X N=40, N2=17 X X X N=40, N2=17 N=40, N2=10, N2=17 N=40, N2=17 N=40, N2=17 N=40, N2=17 N=40, N2=10, N2=17 N=40, N2=10, | |
| N1=40, N2=17 | |
| Fold Change: 2.41 Fold Change: 2.41 Fold Change: 2.41 Fold Change: 2.41 Fold Change: 2.41 Fold Change: 2.56 Fold Change: 2.57 Fold | |
| P-value: .00028 | |
| 113 546 AI659514 110.57+4-163.5 295.11+2422 N1=40, N2=17 | |
| 295.114-242 N1=40, N2=17 Rold Change: 2.56 P-value: .00744 114 562 A1658925 \$42.56+3-47.67 229.65+1-61.58 N1=40, N2=17 Rold Change: 2.07 P-value: .00351 115 565 A1659418 261.02+6-116.11 133.75+1-08.49 N1=40, N2=17 Rold Change: 2.41 P-value: .00368 116 566 A1659533 \$563.44+201.34 291.04+6-136.51 N1=40, N2=17 Rold Change: 2.41 P-value: .00268 117 588 A1689541 \$10.08+20.12 118 589 A1689541 \$10.08+20.12 119 S88 A1680541 \$10.08+20.12 110 S88 A1680541 \$10.08+20.12 110 S88 A1680541 \$10.08+20.12 111 S88 A1680541 \$10.08+20.12 112 S88 A1680541 \$10.08+20.12 113 S89 A1680541 \$10.08+20.12 114 S89 A1680541 \$10.08+20.12 115 S80 A1680541 \$10.08+20.12 116 S80 A1680541 \$10.08+20.12 117 S80 A1680541 \$10.08+20.12 118 S91 A1683911 \$24.46+20.089 \$24.46+20.289 | |
| Ni = 40, N2=17 | |
| Rold Change: 2.56 P-value: 0.0744 | |
| P-value: .00744 | |
| 114 562 | |
| 259,65+1-161.58 N1=40,N2=17 Fold Change: 2.07 Fold Change: 2.41 Fold Change: 2.44 Fold Change: 2.45 Fold Change: 2.45 Fold Change: 2.45 Fold Change: 2.45 Fold Change: 2.45 Fold Change: 2.45 Fold Change: 2.45 Fold Change: 2.44 Fold Change: 2.45 Fold Change: 2.44 Fo | |
| Ni=40, N2=17 | |
| Fold Change: 2.07 Pevalue: .00351 115 | |
| P-value: .00351 | |
| 115 565 AI659418 261,02+116,11 133,75+1,08,49 N1=40, N2=17 X X N1=40, N2=17 N1=40, N2=17 N1=40, N2=17 N1=40, N2=17 N1=40, N2=17 N1=40, N2=17 N2=17 N1=40, N2=17 | |
| 133.75+/108.49 N =40, N2=17 X X X N =40, N2=17 X N = | |
| Ni=40, N2=17 | |
| Fold Change: 2.4 | |
| P-value: .00088 | |
| 116 566 AI659533 553.4+/-201.34 291.04+/-136.51 N1=40, N2=17 X X N1=40, N2=17 X Pold Change: 2.1 P-value: .00023 510.08+/-201.29 106.49+/-4.75 N1=40, N2=17 X N1=40, N2=17 X N1=40, N2=17 Fold Change: 4.54 P-value: .00246 P- | |
| 291,04+/-136.51 N1=40, N2=17 Fold Change; 2.1 P-value: .00023 117 588 A1680541 510.08+/-201.29 510.08+/-201.29 186,08+/-102.82 106,49+4.75 N1=40, N2=17 X N1=40, N2=17 Fold Change; 2.84 P-value: .00266 P-value: .00266 118 591 A1683911 241.46+/-200.89 241.46+/-200.89 | |
| N1=40, N2=17 | |
| Fold Change: 2.1 P-value: .0023 | |
| P-value: .00023 | |
| 117 588 AI680541 510.08+/-201.29 510.08+/-201.29 186.08+/-102.82 106.49+/-44.75 N1=40, N2=17 X N1=40, N2=17 Fold Change: 2.84 Fold Change: 4.54 P-value: 0 118 591 AI683911 241.46+/-200.89 241.46+/-200.89 | |
| 186.084/-102.82 106.491-44.75 186.084/-102.82 106.491-44.75 187.082-17 X 187.082-184 Fold Change: 4.54 Fold | 0 |
| N1=40, N2=17 X N1=40, N2=17 | |
| Fold Changer 2.84 Fold Changer 4.54 P-value: .00246 P-value: .00246 Fold Changer 4.54 P-value: .00246 Fold Changer 4.54 P-value: .00246 P-value: .00246 Fold Changer 4.54 P-value: .00246 P-va | |
| P-value: 0 P-value: 00246 118 591 AI683911 241.46+/-200.89 241.46+/-200.89 | |
| 118 591 AI683911 241.46+/-200.89 241.46+/-200.89 | |
| | |
| 27.24+/-52.93 32.69+/-57.65 | |
| N1=40, N2=17 X N1=40, N2=17 | |
| Fold Change: 4.58 Fold Change: 3.74 | |
| P-value: 0 P-value: .01617 | |
| 119 592 AI684457 96.99+/-74.31 | |
| 253.71+/-245.09 | |
| N1=40, N2=17 X X | |
| Fold Change: 2.25 | |
| P-value: .00425 | |
| 120 593 AI686114 374.48+/-274.59 374.48+/-274.59 | 9 |
| 120.83+/-92.86 76.06+/-83.42 | |
| N1=40, N2=17 X N1=40, N2=17 | |
| Fold Change: 3.03 Fold Change: 4.43 | |
| P-value: .04695 | |
| 121 612 AI701034 215.78+/-96.65 | |
| 111.85+/-71.77 | |
| N1=40, N2=17 X X | |
| Fold Change: 2.11 | |
| P-value: .00036 | |

| # | Seq ID | Genbank | Normal vs All | Normal vs Malignant | Normal vs SII and SIII |
|-----|--------|------------|-------------------|---------------------|------------------------|
| 122 | 618 | AI732274 | 947.08+/-989.69 | | |
| | | | 285.99+/-458.46 | | |
| | | | N1=40, N2=17 | · X | X |
| | | | Fold Change: 3.94 | | |
| | | | P-value: .00251 | | |
| 123 | 619 | AI733679 | 325.9+/-596.22 | | |
| 145 | UL) | ART 100017 | 48.5+/-33.81 | | |
| | | | N1=40, N2=17 | X | x |
| | | | Fold Change: 3.21 | | |
| | | | P-value; .00002 | | |
| 124 | 623 | AI740621 | 231.84+/-247.13 | | |
| 144 | 023 | A1740021 | 77.35+/-124.9 | | |
| | | | N1=40, N2=17 | x | X |
| | | | Fold Change: 2.62 | Λ. | A. |
| | | | P-value: .00315 | | |
| | | 177/2002 | 111.78+/-132.43 | | 111.78+/-132.43 |
| 125 | 627 | AI742002 | | | 388+/-292.79 |
| | | | 379.6+/-168.26 | x | N1=40, N2=17 |
| | | | N1=40, N2=17 | X | |
| | | | Fold Change: 4.61 | | Fold Change: 4.32 |
| | | | P-value: 0 | | P-value: .0111 |
| 126 | 629 | AI742239 | 159.76+/-199.32 | | |
| | | | 419.47+/-377.4 | | |
| | | | N1=40, N2=17 | x | X |
| | | | Fold Change: 3.29 | | |
| | | | P-value: .00013 | | |
| 127 | 631 | AI742490 | 601.57+/-252.84 | | |
| | | | 285.13+/-140.07 | | |
| | | | N1=40, N2=17 | X | x |
| | | | Fold Change: 2.05 | | |
| | | | P-value: .00003 | | |
| 128 | 632 | AI742521 | 215.93+/-234.91 | | 215.93+/-234.91 |
| | | | 23.91+/-22.33 | | 23.3+/-12.66 |
| | | | N1=40, N2=17 | x | N1=40, N2=17 |
| | | | Fold Change: 4.4 | | Fold Change: 4.76 |
| | | | P-value: 0 | | P-value: .00002 |
| 129 | 635 | AI743671 | 582.82+/-317.91 | | |
| | 000 | | 281.49+/-185.49 | | |
| | | | N1=40, N2=17 | x | х |
| | | | Fold Change: 2.26 | | |
| | | | P-value: .00964 | | |
| 130 | 636 | AI743715 | 312.02+/-238.55 | | |
| 130 | 030 | AL/45/15 | 99.48+/-141.4 | | |
| | | | N1=40, N2=17 | х | х |
| | | | | | 24 |
| | | | Fold Change: 3.47 | | |
| | | 17510005 | P-value: .0005 | | |
| 131 | 637 | AI743925 | 663.58+/-309.38 | | |
| | | | 221.31+/-142.28 | | x |
| | | | N1=40, N2=17 | X | X |
| | | | Fold Change: 3.13 | | |
| | | | P-value: 0 | | |
| 132 | 641 | AI751438 | 144.67+/-188.73 | | 144.67+/-188.73 |
| | | | 551.05+/-364.64 | | 612.92+/-347.94 |
| | | | N1=40, N2=17 | · x | N1=40, N2=17 |
| | | | Fold Change: 4.85 | | Fold Change: 5.61 |
| | | | roid Change, 4.65 | | P-value: .02877 |

| # | Seq ID | Genbank | Normal vs All | Normal vs Malignant | Normal vs SII and SIII |
|------|--------|-------------|-------------------|---------------------|------------------------|
| 133 | 643 | AI758223 | 833.52+/-665.83 | | 833.52+/-665.83 |
| | | | 89.52+/-74.43 | | 98.81+/-90.66 |
| l | | | N1=40, N2=17 | x | N1=40, N2=17 |
| | | | Fold Change: 8.3 | | Fold Change: 8 |
| | | | P-value: 0 | | P-value: .02464 |
| 134 | 649 | A1761241 | 883.3+/-332.12 | | |
| ~ . | 0.15 | 121101211 | 415,64+/-208,2 | | |
| ŀ | | | N1=40, N2=17 | x | x |
| | | | Fold Change: 2.21 | | |
| | | | P-value: .00005 | | |
| 135 | 650 | AI761274 | 342.36+/-182.65 | | 342.36+/-182.65 |
| 155 | 050 | 7470127-1 | 121.18+/-64.61 | | 75.25+/-39.87 |
| | | | N1=40, N2=17 | x | N1=40, N2=17 |
| 1 | | | Fold Change: 2.86 | | Fold Change: 4.5 |
| | | | P-value: .00001 | | P-value: .01949 |
| 136 | 652 | AI761844 | 278.83+/-138.41 | | 278.83+/-138.41 |
| 130 | 034 | 231/01044 | 99.54+/-56.16 | | 87.16+/-56.51 |
| l | | | N1=40, N2=17 | x | N1=40, N2=17 |
| | | | Fold Change: 2.79 | | Fold Change: 3.1 |
| | | | P-value: .00001 | | P-value: .02791 |
| 137 | 653 | AI763136 | 282,1+/-149.81 | | |
| 137 | 033 | ALTOSISO | 118.7+/-131.83 | | |
| 1 | | | N1=40, N2=17 | x | x |
| | | | Fold Change: 2.53 | | |
| | | | P-value: .00163 | | |
| 138 | 655 | AI766029 | 271.74+/-528.19 | | 271.74+/-528.19 |
| 130 | 055 | A1700029 | 22.11+/-18.39 | | 30,31+/-29,22 |
| | | | N1=40, N2=17 | x | N1=40, N2=17 |
| 1 | | | Fold Change: 3.71 | 1. | Fold Change: 3.07 |
| | | | P-value: 0 | | P-value: .01978 |
| 139 | 657 | AI768325 | 114.7+/-66.43 | | |
| 137 | 057 | AL 100325 | 257.51+/-172.22 | | |
| 1 | | | N1=40, N2=17 | X | x |
| | | | Fold Change: 2.12 | | |
| | | | P-value: .00044 | | |
| 140 | 664 | AI791182 | 286,48+/-162,61 | | |
| 140 | 004 | A1771102 | 621.07+/-388.18 | | |
| 1 | | | N1=40, N2=17 | x | x |
| 1 | | | Fold Change: 2.07 | ,, | |
| 1 | | | P-value: .00052 | | |
| 141 | 668 | AI792635 | 1-14440, 100002 | | 800.24+/-717.81 |
| 141 | 000 | Par. / 2033 | | | 1968.88+/-866 |
| J | | | x | x | N1=40, N2=17 |
| | | | Α. | | Fold Change: 4.27 |
| | | | | | P-value: .0038 |
| 142 | 674 | AI797276 | 271.48+/-136.73 | | 271.48+/-136.73 |
| 142 | 074 | AL17/2/0 | 106.25+/-58.1 | | 76.49+/-46.61 |
| | | | N1=40, N2=17 | x | N1=40, N2=17 |
| 1 | | | Fold Change: 2.56 | 46 | Fold Change: 3.58 |
| | | | P-value: .00001 | | P-value: .02759 |
| 1.12 | C#0 | AI799784 | 603.99+/-383.42 | | 603.99+/-383.42 |
| 143 | 678 | AL /99/84 | 93.05+/-88.68 | | 82.71+/-78.29 |
| 1 | | | | x | N1=40, N2=17 |
| | | | N1=40, N2=17 | Α. | Fold Change: 7.34 |
| | | | Fold Change: 6.66 | | P-value: .01379 |
| L | | | P-value: 0 | | r-value: .013/9 |

| # | Seq ID | Genbank | Normal vs Ali | Normal vs Malignant | Normal vs SII and SIII |
|-----|--------|----------|------------------------------------|---------------------|---------------------------------|
| 144 | 684 | AI804054 | 302.97+/-234.41 | | 302.97+/-234.41 |
| 1 | | | 108.18+/-91.04 | | 77.58+/-36.68 |
| 1 | | | N1=40, N2=17 | x | N1=40, N2=17 |
| 1 | | | Fold Change: 2.83 | | Fold Change: 3,3 |
| 1 | | | P-value: .00011 | | P-value: .01862 |
| 145 | 687 | AI806324 | 211.46+/-131.17 | | |
| 1 | | | 108.84+/-79,43 | | |
| 1 | | | N1=40, N2=17 | x | X |
| 1 | | | Fold Change: 2.03 | | |
| | | | P-value: .00874 | | |
| 146 | 691 | AI809953 | 383.43+/-189.32 | | |
| ļ | | | 120,52+/-100,18 | | |
| | | | N1=40, N2=17 | x | X |
| 1 | | | Fold Change: 3.27 | | |
| | | | P-value: .00013 | | |
| 147 | 693 | AI810266 | 68.88+/-106.64 | | |
| 1 | | | 761.49+/-1126.65 | | |
| 1 | | | N1=40, N2=17 | X | x |
| } | | | Fold Change: 6.3 | | |
| | | | P-value: .00013 | | |
| 148 | 694 | AI810764 | 202.16+/-159.83 | | |
| 1 | | | 1084.09+/-1401.59 | | |
| 1 | | | N1=40, N2=17 | X | x |
| 1 | | | Fold Change: 4.41 | | |
| | | | P-value: .00007 | | |
| 149 | 701 | AI816835 | 360.85+/-289.77 | | |
| 1 | | | 171.05+/-158.66 | | |
| { | | | N1=40, N2=17 | x | x |
| ł | | | Fold Change: 2.13 | | |
| | | | P-value: .00229 | | |
| 150 | 704 | AI817967 | | | 112.71+/-118.41 |
| Į | | | | | 308.86+/-160.78 |
| 1 | | | X | X | N1=40, N2=17 |
| l | | | | | Fold Change: 3.45 |
| | | | | | P-value: .00951 |
| 151 | 706 | AI818579 | 394.08+/-228.07 | | |
| 1 | | | 204.91+/-197.94 | *** | w |
| 1 | | | N1=40, N2=17 | X | X |
| | | | Fold Change: 2.13 | | |
| 152 | 712 | AI821472 | P-value: .00391 | | 519.11+/-694.13 |
| 152 | 114 | A18214/2 | 519.11+/-694.13 -5.59+/-218.89 | | -49.74+/-70.96 |
| l | | | | • | |
| 1 | | | N1=40, N2=17 | X | N1=40, N2=17 |
| 1 | | | Fold Change: 5.69 | | Fold Change: 9.33 P-value: 0 |
| 153 | 713 | AI823572 | P-value: .00005 232.21+/-195.63 | | r-value; o |
| 133 | /13 | A10435/4 | 91.57+/-60.62 | | |
| į | | | N1=40, N2=17 | x | x |
| } | | | Fold Change: 2.43 | ^ | ^ |
| | | | P-value: .00008 | | |
| 154 | 721 | AI825936 | 229.86+/-148.12 | | |
| 134 | 141 | A1043930 | 98.58+/-81.47 | | |
| 1 | | | 98.58+7-81.47 N1=40, N2=17 | х | x |
| 1 | | | Fold Change: 2.58 | Λ | ^ |
| } | | | P-value: .00016 | | |
| | | | r-vame: .00016 | | |

| # | Seq ID | Genbank | Normal vs All | Normal vs Malignant | Normal vs SII and SIII |
|-----|--------|-----------|-------------------|---------------------|------------------------|
| 155 | 722 | A1826437 | 45.86+/-118.99 | | |
| 100 | | | 281.35+/-448.25 | | |
| | | | N1=40, N2=17 | x | x |
| | | | Fold Change: 3.03 | | |
| | | | P-value: .0122 | | |
| 156 | 744 | AI863167 | 183.76+/-73.48 | | |
| 130 | / | 711005107 | 406.96+/-190.24 | | |
| | | | N1=40, N2=17 | x | x |
| | | | Fold Change: 2.16 | | |
| | | | P-value: 0 | | |
| 157 | 747 | AI864898 | 401.86+/-258.51 | | |
| 13/ | /4/ | A1004020 | 75,46+/-68.5 | | |
| | | | N1=40, N2=17 | x | x |
| | | | Fold Change: 5.61 | | |
| | | | P-value: 0 | | |
| 158 | 750 | AI871044 | 766.39+/-500.99 | | 766.39+/-500.99 |
| 138 | 150 | A10/1044 | 189.5+/-179.55 | | 84.85+/-70.19 |
| | | | N1=40, N2=17 | x | N1=40, N2=17 |
| | | | Fold Change: 4.03 | | Fold Change: 8.12 |
| | | | P-value: .00001 | | P-value: .00884 |
| 159 | 751 | AI872267 | 267.23+/-203.1 | | A 1000001 |
| 159 | /51 | A10/220/ | 627.26+/-368.25 | | |
| | | | N1=40, N2=17 | x | x |
| | | | Fold Change: 2.55 | A . | |
| | | | P-value: .00015 | | |
| 160 | 752 | AI879337 | 431.51+/-184.18 | | |
| 100 | 152 | A10/933/ | 215.5+/-115.86 | | |
| | | | N1=40, N2=17 | x | x |
| | | | Fold Change: 2.18 | | |
| | | | P-value: .00062 | | |
| 161 | 758 | AI888322 | I Made Toolean | | 319.22+/-320.74 |
| 101 | 150 | A1000322 | | | 71.54+/-51.15 |
| | | | x | х | N1=40, N2=17 |
| | | | ** | | Fold Change: 3.78 |
| | | | | | P-value: .03277 |
| 162 | 772 | AI916544 | 151.27+/-163.24 | | |
| 202 | ,,,, | | 373.43+/-334.2 | | |
| | | | N1=40, N2=17 | x | x |
| | | | Fold Change: 2.45 | | |
| | | | P-value: .00524 | | |
| 163 | 775 | AI917901 | 601.53+/-812.45 | | 601.53+/-812.45 |
| 130 | | | 76.98+/-131.25 | | 26.66+/-20.01 |
| | | | N1=40, N2=17 | x | N1=40, N2=17 |
| | | | Fold Change: 4.95 | *- | Fold Change: 7.3 |
| | | | P-value: .00005 | | P-value: .00001 |
| 164 | 780 | AI924465 | 448.27+/-478.27 | | |
| 104 | 700 | .11/14/00 | 149.48+/-115.97 | | |
| | | | N1=40, N2=17 | x | x |
| | | | Fold Change: 2.43 | ** | |
| | | | P-value: .00214 | | |
| 165 | 787 | AI934361 | 220.01+/-243.16 | | 220.01+/-243.16 |
| 105 | /6/ | A1934301 | 54.43+/-44.52 | | 52.02+/-37.1 |
| | | | N1=40, N2=17 | x | N1=40, N2=17 |
| | | | Fold Change: 3.1 | Λ. | Fold Change: 3.01 |
| | | | P-value: .00001 | | P-value: .03711 |
| | | | r-value: .00001 | | 1-7414005/11 |

| # | Seq ID | Genbank | Normal vs All | Normal vs Malignant | Normal vs SII and SIII |
|-----|--------|--------------|-------------------|---------------------|------------------------|
| 166 | 789 | AI934881 | 316.72+/-226.37 | | |
| | | | 659.59+/-486.96 | | |
| | | | N1=40, N2=17 | x | x |
| | | | Fold Change: 2,01 | | |
| | | | P-value: .00378 | | |
| 167 | 816 | AI968151 | 127.39+/-61.78 | | |
| | | | 376.92+/-292.97 | | |
| | | | N1=40, N2=17 | X | x |
| | | | Fold Change: 2.53 | | |
| | | | P-value: .00031 | | |
| 168 | 817 | AI968379 | 295.46+/-388.02 | | 295.46+/-388.02 |
| 200 | | | -8.49+/-25.52 | | .59÷/-34.6 |
| | | | N1=40, N2=17 | X | N1=40, N2=17 |
| | | | Fold Change: 6.27 | | Fold Change: 5.43 |
| | | | P-value: 0 | | P-value: .00032 |
| 169 | 818 | AI968904 | 738.79+/-292.65 | | |
| | | | 307.62+/-119.37 | | |
| | | | N1=40, N2=17 | x | x |
| | | | Fold Change: 2.35 | | |
| i | | | P-value: 0 | | |
| 170 | 830 | AI972498 | 286.51+/-112.64 | | |
| 1,0 | 050 | 1220 /2 10 4 | 135.46+/-66.44 | | |
| | | | N1=40, N2=17 | x | x |
| | | | Fold Change: 2.18 | | |
| | | | P-value: .00003 | | |
| 171 | 832 | AI972873 | 436.16+/-215 | | |
| | | | 132.01+/-99.1 | | |
| | | | N1=40, N2=17 | x | x |
| | | | Fold Change: 3.85 | | |
| | | | P-value: 0 | _ | |
| 172 | 838 | AI983045 | 281.02+/-338.08 | | 281.02+/-338.08 |
| | | | 40.45+/-125.75 | | -9.19+/-15.96 |
| ŀ | | | N1=40, N2=17 | x | N1=40, N2=17 |
| | | | Fold Change: 4.78 | | Fold Change: 7.52 |
| | | | P-value: 0 | | P-value: 0 |
| 173 | 857 | AL037805 | | | 614.2+/-317.15 |
| 1 | | | | | 183.89+/-87.99 |
| 1 | | | x | X | N1=40, N2=17 |
| | | | | | Fold Change: 3.13 |
| | | | | | P-value: .01435 |
| 174 | 865 | AL040912 | 304.56+/-132.78 | | |
| | | | 112.19+/-70.33 | | ** |
| | | | N1=40, N2=17 | X | X |
| | | | Fold Change: 2.8 | | |
| | | | P-value: .00006 | | |
| 175 | 867 | AL042492 | 809.69+/-853.09 | | 1022.91+/-907.07 |
| 1 | | | 72.75+/-93.44 | ** | 85.76+/-67.41 |
| 1 | | | N1=40, N2=17 | · x | N1=40, N2=17 |
| 1 | | | Fold Change: 9.48 | | Fold Change: 8.09 |
| | | | P-value: 0 | | P-value: .00176 |
| 176 | 876 | AL046941 | 428.58+/-238.89 | | 428.58+/-238.89 |
| | | | 146.79+/-176.57 | | 55.32+/-48.1 |
| | | | N1=40, N2=17 | X | N1=40, N2=17 |
| 1 | | | Fold Change: 4.06 | | Fold Change: 7.58 |
| 1 | | | P-value: .00007 | _ | P-value: .01267 |

| # | Seq ID | Genbank | Normal vs All | Normal vs Malignant | Normal vs SII and SIII |
|-----|--------|-------------|-------------------|---------------------|------------------------|
| 177 | 881 | AL048962 | 944+/-354.29 | | 944+/-354.29 |
| | | | 399.3+/-211.63 | | 289.62+/-184.81 |
| | | | N1=40, N2=17 | x | N1=40, N2=17 |
| | | | Fold Change: 2.5 | | Fold Change: 3.52 |
| | | | P-value: .00001 | | P-value: .03411 |
| 178 | 893 | AL050367 | 257.59+/-77.75 | | 257.59+/-77.75 |
| 170 | 050 | 122000007 | 111.77+/-59.21 | | 76.12+/-36.74 |
| | | | N1=40, N2=17 | x | N1=40, N2=17 |
| | | | Fold Change: 2.47 | | Fold Change: 3.45 |
| | | | P-value: 0 | | P-value: .01201 |
| 179 | 894 | AL079279 | 313,49+/-189,76 | | |
| 115 | 024 | ALGINETY | 127.56+/-77.14 | | |
| | | | N1=40, N2=17 | x | x |
| | | | Fold Change: 2.4 | | == |
| | | | P-value: .00036 | | |
| 100 | 896 | AL079707 | 261.69+/-226.08 | | |
| 180 | 896 | ALU/9/0/ | | | |
| | | | 73.98+/-35.27 | x | x |
| | | | N1=40, N2=17 | ^ | |
| | | | Fold Change: 3.16 | | |
| | | | P-value: 0 | | 234.63+/-113.05 |
| 181 | 902 | AL118746 | 234.63+/-113.05 | | 46.19+/-34.43 |
| | | | 84.77+/-47.29 | 37 | N1=40, N2=17 |
| | | | N1=40, N2=17 | x | |
| | | | Fold Change: 2.86 | | Fold Change: 5.14 |
| | | | P-value: .00001 | | P-value: .0179 |
| 182 | 905 | AW000952 | 98.9+/-72.25 | | |
| | | | 204.67+/-105.21 | | |
| | | | N1=40, N2=17 | X | X |
| | | | Fold Change: 2.16 | | |
| | | | P-value: .00011 | | |
| 183 | 907 | AW002846 | 283.14+/-201.6 | | |
| | | | 119.62+/-87.38 | | |
| | | | N1=40, N2=17 | X | x |
| | | | Fold Change: 2.43 | | |
| | | | P-value: .00065 | | |
| 184 | 908 | AW002941 | 959.64+/-342.08 | | |
| | | | 493.25+/-243.21 | | |
| | | | N1=40, N2=17 | x | X |
| | | | Fold Change: 2.13 | | |
| | | | P-value: .0001 | | |
| 185 | 916 | AW006235 | 346.9+/-210.26 | | |
| _00 | 210 | | 121.01+/-58.03 | | |
| | | | N1=40, N2=17 | x | x |
| | | | Fold Change: 2.69 | | |
| | | | P-value: 0 | | |
| 186 | 917 | AW006352 | 235,29+/-179.11 | | |
| 100 | J1 1 | A 11 000334 | 534.97+/-420.56 | | |
| | | | N1=40, N2=17 | x | x |
| | | | Fold Change: 2.17 | ** | |
| | | | P-value: .00953 | | |
| 105 | 001 | 1337007000 | 223.2+/-116.87 | | 223.2+/-116.87 |
| 187 | 921 | AW007080 | | | 36.39+/-14.01 |
| | | | 69.24+/-50.48 | х | N1=40, N2=17 |
| | | | N1=40, N2=17 | A | Fold Change: 5.16 |
| | | | Fold Change: 3.27 | | P-value: .0001 |
| | | | P-value: .00001 | | r-value: .0001 |

| # | Seq ID | Genbank | Normal vs All | Normal vs Malignant | Normal vs SII and SIII |
|-----|--------|------------|-------------------|---------------------|------------------------|
| 188 | 926 | AW007803 | 153.39+/-142.06 | | |
| | | | 442.5+/-397.54 | | |
| | | | N1=40, N2=17 | x | x |
| | | | Fold Change: 2,55 | | |
| | | | P-value: .00867 | | |
| 189 | 931 | AW014155 | 214.48+/-209.56 | | |
| | | | 624.36+/-372.34 | | |
| | | | N1=40, N2=17 | x | X |
| | | | Fold Change: 3.15 | | |
| | | | P-value: .00005 | | |
| 190 | 953 | AW051492 | 442,65+/-332,99 | | |
| 1,0 | ,555 | 1411051452 | 203.39+/-140.38 | | |
| | | | N1=40, N2=17 | x | x |
| | | | Fold Change: 2.22 | | |
| | | | P-value: .00151 | | |
| 191 | 957 | C17781 | 229.36+/-141.71 | | |
| 191 | 951 | C1//01 | 84.23+/-69.19 | | |
| | | | N1=40, N2=17 | x | x |
| | | | Fold Change: 2.59 | A | |
| | | | P-value: .00012 | | |
| 192 | 975 | F22640 | 416.82+/-153.5 | | |
| 192 | 9/5 | F22040 | 204.94+/-169.19 | | |
| | | | | x | x |
| | | | N1=40, N2=17 | ^ | |
| | | | Fold Change: 2.37 | | |
| | | WW- 4840 | P-value: .00007 | | 288.53+/-212.27 |
| 193 | 985 | H16568 | 288.53+/-212.27 | | 32.47+/-46.57 |
| | | | 74.99+/-76.74 | x | N1=40, N2=17 |
| | | | N1=40, N2=17 | | Fold Change: 5.01 |
| | | | Fold Change: 3.32 | | P-value: .01332 |
| | | | P-value: .00019 | | F-value: .01332 |
| 194 | 988 | H30384 | 194.93+/-133.51 | | |
| | | | 479.18+/-480.95 | ** | x |
| | | | N1=40, N2=17 | x | X |
| | | | Fold Change: 2.18 | | |
| | | | P-value: .00329 | | |
| 195 | 992 | H54254 | 377.04+/-687.01 | | 377.04+/-687.01 |
| | | | 38.27+/-23.01 | | 36.82+/-32.95 |
| | | | N1=40, N2=17 | X | N1=40, N2=17 |
| | | | Fold Change: 4.25 | | Fold Change: 4.51 |
| | | | P-value: 0 | | P-value: .00966 |
| 196 | 997 | H92988 | 390.91+/-149.13 | | |
| | | | 205.04+/-140.06 | | |
| | | | N1=40, N2=17 | x | X |
| | | | Fold Change: 2.33 | | |
| | | | P-value: .00168 | | |
| 197 | 1074 | N42752 | 63.77+/-48.02 | | |
| | | | 291.54+/-224.99 | | |
| | | | N1=40, N2=17 | x | X |
| | | | Fold Change: 3.86 | | |
| | | | P-value: .00006 | | |
| 198 | 1085 | N56877 | 109,5+/-80,79 | | |
| 1/0 | 1000 | 1150077 | 402.12+/-388.61 | | |
| | | | N1=40, N2=17 | x | x |
| | | | Fold Change: 3 | ** | |
| | | | P-value: .00087 | | |

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| # | Seq ID | Genbank | Normal vs All | Normal vs Malignant | Normal vs SII and SIII |
|-----|--------|---------|-------------------|---------------------|------------------------|
| 199 | 1090 | N63913 | 458.01+/-316.71 | | 458.01+/-316.71 |
| | | | 67.39+/-79.5 | | 8.79+/-40.55 |
| | | | N1=40, N2=17 | x | N1=40, N2=17 |
| | | | Fold Change: 6.57 | - | Fold Change: 11.59 |
| | | | P-value: 0 | | P-value: .00004 |
| 200 | 1101 | R08000 | 502.76+/-694.51 | | 502.76+/-694.51 |
| 200 | 1101 | 200000 | 82.03+/-53.39 | | 90.53+/-93.25 |
| | | | N1=40, N2=17 | x | N1=40, N2=17 |
| | | | Fold Change: 3.78 | | Fold Change: 4.05 |
| | | | P-value: 0 | | P-value: .04964 |
| 201 | 1104 | R20784 | 1112.78+/-843.96 | | |
| 201 | 110-7 | 1120704 | 359,34+/-233,36 | | |
| | | | N1=40, N2=17 | x | x |
| | | | Fold Change: 2.91 | 24 | |
| | | | P-value: .00005 | | |
| 202 | 1105 | R39938 | 111.89+/-67.41 | | |
| 202 | 1105 | K39936 | 222.22+/-111.16 | | |
| | | | N1=40, N2=17 | X | х . |
| | | | Fold Change: 2.12 | Α. | |
| | | | | | |
| | | | P-value: .00002 | | |
| 203 | 1106 | R42575 | 90.17+/-38.15 | | |
| | | | 215.36+/-156.29 | х | x |
| | | | N1=40, N2=17 | X | х |
| | | | Fold Change: 2.01 | | |
| | | | P-value: .00211 | | 200 25: 1400 05 |
| 204 | 1112 | R54660 | 200.26+/-133.86 | | 200.26+/-133.86 |
| | | | 48.69+/-33.36 | | 29.39+/-27.33 |
| | | | N1=40, N2=17 | Х | N1=40, N2=17 |
| | | | Fold Change: 3.43 | | Fold Change: 4.69 |
| | | | P-value: 0 | | P-value: .0025 |
| 205 | 1116 | R70255 | 241.29+/-181.34 | | 241.29+/-181.34 |
| | | | 14.29+/-38.71 | | -8.34+/-15.47 |
| | | | N1=40, N2=17 | x | N1=40, N2=17 |
| | | | Fold Change: 5.79 | | Fold Change: 7.98 |
| | | | P-value: 0 | | P-value: 0 |
| 206 | 1118 | R74561 | 425.23+/-350.96 | | |
| | | | 879,43+/-654.71 | | |
| | | | N1=40, N2=17 | x | x |
| | | | Fold Change: 2.16 | | |
| | | | P-value: .0019 | | |
| 207 | 1119 | R83604 | 304.76+/-867.74 | | |
| 207 | 1117 | 200000 | -32.63+/-64.18 | | |
| | | | N1=40, N2=17 | x | x |
| | | | Fold Change: 3.15 | | |
| | | | P-value: .00017 | | |
| 208 | 1125 | T61106 | 180.38+/-114.3 | | |
| 200 | 1145 | 101100 | 349.03+/-164.74 | | |
| | | | N1=40, N2=17 | x | x |
| | | | Fold Change: 2.35 | Α. | Α. |
| | | | | | |
| | | | P-value; .00001 | | 166.2+/-116.99 |
| 209 | 1132 | T85314 | | | 644.58+/-401.95 |
| | | | | ** | |
| | | | X | X | N1=40, N2=17 |
| | | | | | Fold Change: 4.09 |
| | | | | | P-value: .03546 |

| # | Seq ID | Genbank | Normal vs All | Normal vs Malignant | Normal vs SII and SIII |
|----------|--------|-----------|-------------------|---------------------|------------------------|
| 210 | 1171 | W02823 | 217.4+/-87.18 | | 217.4+/-87.18 |
| | | | 81.39+/-47.18 | | 53.69+/-25.71 |
| | | | N1=40, N2=17 | X | N1=40, N2=17 |
| | | | Fold Change: 2.83 | | Fold Change: 3.92 |
| | | | P-value: .00001 | | P-value: .00806 |
| 211 1173 | 1173 | W07043 | 299.21+/-164.12 | | 299.21+/-164.12 |
| | | | 105.66+/-83.76 | | 59.94+/-40.54 |
| | | | N1=40, N2=17 | x | NI=40, N2=17 |
| | | | Fold Change: 2.82 | | Fold Change: 4.46 |
| | | | P-value: .00008 | | P-value: .01951 |
| 212 | 1174 | W07304 | 1139.71+/-444.58 | | 1139.71+/-444.58 |
| | | | 502.93+/-458.99 | | 349.93+/-213.71 |
| | | | N1=40, N2=17 | x | N1=40, N2=17 |
| | | | Fold Change: 2.64 | | Fold Change: 3.49 |
| | | | P-value: .00012 | | P-value: .04978 |
| 213 | 1180 | W27541 | 1 /44001100012 | | 486.94+/-189.31 |
| 213 | 1100 | 112/541 | | | 113.57+/-41.71 |
| | | | х . | x | N1=40, N2=17 |
| | | | Α. | ^ | Fold Change: 4.17 |
| | | | | | P-value: .0025 |
| 214 | 1183 | W32480 | 720.17+/-951.89 | | 720.17+/-951.89 |
| 214 | 1183 | W32480 | 76.05+/-158.18 | | 18.91+/-12.14 |
| | | | N1=40, N2=17 | x | N1=40, N2=17 |
| | | | | | Fold Change: 12.97 |
| | | | Fold Change: 7.94 | | P-value: 0 |
| | | **** | P-value: 0 | | P-value; 0 |
| 215 | 1184 | W37770 | 208.87+/-62 | | |
| | | | 108.93+/-55.29 | x | x |
| | | | N1=40, N2=17 | X | X |
| | | | Fold Change: 2.1 | | |
| | | | P-value: .00006 | | |
| 216 | 1185 | W37896 | 499.73+/-192.2 | | |
| | | | 1636.96+/-1336.48 | | , ,, |
| | | | N1=40, N2=17 | X | , X |
| | | | Fold Change: 2.49 | | |
| | | | P-value: .00074 | | |
| 217 | 1198 | W72338 | 464.08+/-121.49 | | |
| | | | 964.48+/-427.69 | | |
| | | | N1=40, N2=17 | X | X |
| | | | Fold Change: 2 | | |
| | | | P-value: 0 | | |
| 218 | 1199 | W72347 | 368.08+/-157.32 | | |
| | | | 134.9+/-113.13 | | |
| | | | N1=40, N2=17 | x | x |
| | | | Fold Change: 3.01 | | |
| | | | P-value: .00008 | | |
| 219 | 1200 | W72407 | 234.77+/-159.7 | | 234.77+/-159.7 |
| | | | 50.76+/-52.77 | | 44,31+/-63.03 |
| | | | N1=40, N2=17 | x | N1=40, N2=17 |
| | | | Fold Change: 4.25 | | Fold Change: 5.12 |
| | | | P-value: 0 | | P-value: .03464 |
| 220 | 1201 | W72511 | 988,5+/-437,53 | | 2 10001001 |
| 220 | 1201 | ** / 2311 | 477.34+/-271.59 | | |
| | | | N1=40, N2=17 | x | х |
| | | | Fold Change: 2.11 | A | A |
| | | | | | |

| # | Seq ID | Genbank | Normal vs All | Normal vs Malignant | Normal vs SII and SIII |
|--------|--------|---------|-------------------|---------------------|------------------------|
| 221 1: | 1204 | W73386 | 248.29+/-403.35 | 469.37+/-905.14 | |
| | | | 35.26+/-68.97 | 101.34+/-51.41 | |
| | | | N1=40, N2=17 | N1=17, N2=7 | x |
| | | | Fold Change: 3.18 | Fold Change: 2.82 | |
| | | | P-value: .0001 | P-value: .01061 | |
| 222 | 1207 | W73890 | 223.01+/-130.77 | | |
| | | | 84.82+/-49.32 | | |
| | | | N1=40, N2=17 | x | X |
| | | | Fold Change: 2.49 | | |
| | | | P-value: 0 | | |
| 223 | 1246 | Z99386 | 611.71+/-209.91 | | |
| | | | 288.23+/-106.96 | | |
| | | | N1=40, N2=17 | x | x |
| | | | Fold Change: 2.19 | | |
| | | | P-value: .00001 | | |

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What is claimed is:

- A method of diagnosing breast cancer in a patient, comprising:
 - (a) detecting the level of expression in a tissue sample of two or more genes from les 1-5; wherein differential expression of the genes in Tables 1-5 is indicative of breas
- Tables 1-5; wherein differential expression of the genes in Tables 1-5 is indicative of breast cancer.
 - 2. A method of detecting the progression of breast cancer in a patient, comprising:
 - (a) detecting the level of expression in a tissue sample of two or more genes from
- 0 Tables 1-5; wherein differential expression of the genes in Tables 1-5 is indicative of breast cancer progression.
 - 3. A method of monitoring the treatment of a patient with breast cancer, comprising:
 - (a) administering a pharmaceutical composition to the patient;
- 15 (b) preparing a gene expression profile from a cell or tissue sample from the patient; and
 - (c) comparing the patient gene expression profile to a gene expression from a cell population selected from the group consisting of normal breast cells and cancerous breast cells.

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- 4. A method of treating a patient with breast cancer, comprising:
- (a) administering to the patient a pharmaceutical composition, wherein the composition alters the expression of at least one gene in Tables 1-5;
- (b) preparing a gene expression profile from a cell or tissue sample from the patient comprising tumor cells; and
 - (c) comparing the patient expression profile to a gene expression profile selected from the group consisting of normal breast cells and cancerous breast cells.
 - A method of typing breast cancer in a patient, comprising:
- (a) detecting the level of expression in a tissue sample of two or more genes from Tables 1-5; wherein differential expression of the genes in Tables 1-5 is indicative of a type of breast cancer selected from a group consisting of infiltrating ductal carcinoma, microinvasive carcinoma, cribiform carcinoma, stage I carcinoma, stage II carcinoma or lobular carcinoma.

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- A method of detecting the presence or progression of infiltrating ductal carcinoma in a patient, comprising:
- (a) detecting the level of expression in a tissue sample of two or more genes from
- Tables 1-5; wherein differential expression of the genes in Tables 1-5 is indicative of infiltrating ductal carcinoma progression.
- A method of monitoring the treatment of a patient with infiltrating ductal carcinoma, comprising:
 - (a) administering a pharmaceutical composition to the patient;

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- (b) preparing a gene expression profile from a cell or tissue sample from the patient; and
- (c) comparing the patient gene expression profile to a gene expression from a cell population comprising normal breast cells or to a gene expression profile from a cell population comprising infiltrating ductal carcinoma cells or to both.
- 8. A method of treating a patient with infiltrating ductal carcinoma, comprising:
- (a) administering to the patient a pharmaceutical composition, wherein the composition alters the expression of at least one gene in Tables 1-5;
- (b) preparing a gene expression profile from a cell or tissue sample from the patient comprising infiltrating ductal carcinoma cells; and
- (c) comparing the patient expression profile to a gene expression profile from an untreated cell population comprising infiltrating ductal carcinoma cells.
- 25 9. A method of diagnosing a microinvasive form of breast tumor in a patient, comprising:
 - (a) detecting the level of expression in a tissue sample of two or more genes from Tables 1-5; wherein differential expression of the genes in Tables 1-5 is indicative of a microinvasive form of breast cancer.
 - 10. A method of detecting the progression of a microinvasive for of breast cancer in a patient, comprising:

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- (a) detecting the level of expression in a tissue sample of two or more genes from Tables 1-5; wherein differential expression of the genes in Tables 1-5 is indicative of the progression of a microinvasive form of breast cancer.
- 5 11. A method of monitoring the treatment of a patient with a microinvasive form of breast cancer, comprising:
 - (a) administering a pharmaceutical composition to the patient;

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- (b) preparing a gene expression profile from a cell or tissue sample from the patient; and
- (c) comparing the patient gene expression profile to a gene expression from a cell population comprising normal breast cells or to a gene expression profile from a cell population comprising microinvasive breast cancer cells or to both.
- A method of treating a patient with a microinvasive form of breast cancer, comprising;
- (a) administering to the patient a pharmaceutical composition, wherein the composition alters the expression of at least one gene in Tables 1-5;
- (b) preparing a gene expression profile from a cell or tissue sample from the patient comprising microinvasive breast cancer cells; and
- (c) comparing the patient expression profile to a gene expression profile from an untreated cell population comprising microinvasive breast cancer cells.
 - 13. A method of differentiating microinvasive breast cancer from a benign growth in a patient, comprising:
- (a) detecting the level of expression in a tissue sample of two or more genes from Tables 1-5; wherein differential expression of the genes in Tables 1-5 is indicative of microinvasive breast cancer rather than benign growth.
- 14. A method of screening for an agent capable of modulating the onset or progression of breast cancer, comprising:
- (a) preparing a first gene expression profile of a cell population comprising breast cancer cells, wherein the expression profile determines the expression level of one or more genes from Tables 1-5;
 - (b) exposing the cell population to the agent;

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- (c) preparing second gene expression profile of the agent-exposed cell population; and
 - (d) comparing the first and second gene expression profiles.
- 5 15. The method of claim 14, wherein the breast cancer is a infiltrating ductal carcinoma.
 - 16. The method of claim 14, wherein the breast cancer is a microinvasive breast cancer.
 - A composition comprising at least two oligonucleotides, wherein each of the oligonucleotides comprises a sequence that specifically hybridizes to a gene in Tables 1-5.
 - A composition according to claim 17, wherein the composition comprises at least 3 oligonucleotides.
- 15 19. A composition according to claim 17, wherein the composition comprises at least 5 oligonucleotides.
 - A composition according to claim 17, wherein the composition comprises at least 7 oligonucleotides.

A composition according to claim 17, wherein the composition comprises at least 10 oligonucleotides.

- A composition according to any one of claims 17-21, wherein the oligonucleotides
 are attached to a solid support.
 - 23. A composition according to claim 22, wherein the solid support is selected from a group consisting of a membrane, a glass support, a filter, a tissue culture dish, a polymeric material, a bead and a silica support.
 - 24. A solid support comprising at least two oligonucleotides, wherein each of the oligonucleotides comprises a sequence that specifically hybridizes to a gene in Tables 1-5.

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- A solid support according to claim 24, wherein the oligonucleotides are covalently attached to the solid support.
- A solid support according to claim 24, wherein the oligonucleotides are noncovalently attached to the solid support.
 - A solid support according to claim 24, wherein the support comprises at least about
 different oligonucleotides in discrete locations per square centimeter.
- 10 28. A solid support according to claim 24, wherein the support comprises at least about 100 different oligonucleotides in discrete locations per square centimeter.
 - A solid support according to claim 24, wherein the support comprises at least about
 1000 different oligonucleotides in discrete locations per square centimeter.
 - A solid support according to claim 24, wherein the support comprises at least about 10,000 different oligonucleotides in discrete locations per square centimeter.
 - 31. A computer system comprising:

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- (a) a database containing information identifying the expression level in breast tissue of a set of genes comprising at least two genes in Tables 1-5; and
 - (b) a user interface to view the information.
- A computer system of claim 31, wherein the database further comprises sequence
 information for the genes.
 - 33. A computer system of claim 31, wherein the database further comprises information identifying the expression level for the genes in normal breast tissue.
- 34. A computer system of claim 31, wherein the database further comprises information identifying the expression level for the genes in breast cancer tissue.
 - A computer system of claim 34, wherein the breast cancer tissue comprises infiltrating ductal carcinoma cells.

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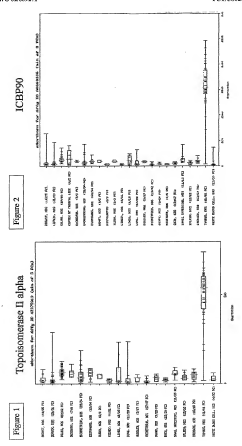
- A computer system of claim 34, wherein the breast cancer tissue comprises microinvasive breast cancer cells.
- 5 37. A computer system of any of claims 31-36, further comprising records including descriptive information from an external database, which information correlates said genes to records in the external database.
 - 38. A computer system of claim 37, wherein the external database is GenBank.

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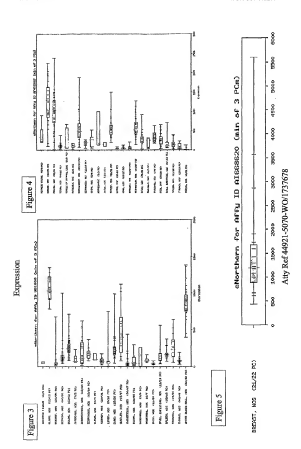
- 39. A method of using a computer system of any one of claims 31-36 to present information identifying the expression level in a tissue or cell of at least one gene in Tables 1-5, comprising:
- (a) comparing the expression level of at least one gene in Tables 1-5 in the tissue or cell to the level of expression of the gene in the database.
 - A method of claim 39, wherein the expression level of at least two genes are compared.
- A method of claim 39, wherein the expression level of at least five genes are compared.
 - A method of claim 39, wherein the expression level of at least ten genes are compared.
 - 43. A method of claim 39, further comprising displaying the level of expression of at least one gene in the tissue or cell sample compared to the expression level in breast cancer.
 - 44. A kit comprising at least one solid support of any one of claims 24-30 packaged with gene expression information for said genes.
 - 45. A kit of claim 44, wherein the gene expression information comprises gene expression levels in a breast cancer tissue or cell sample.

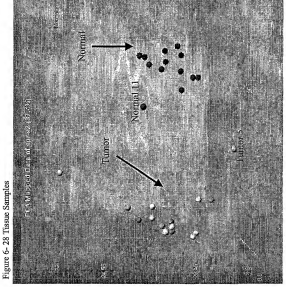
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46. A kit of claim 45, wherein the gene expression information is in an electronic format.

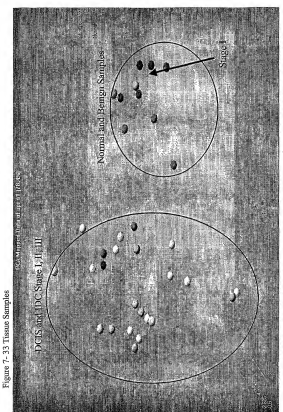


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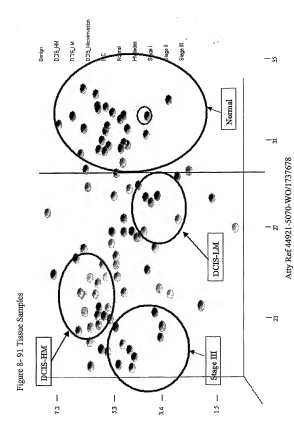




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